Requester

Name: BIANCHI, KRISTIN A

Organization: TC 1600

Art Unit: 1622 Employee Number:

Office Location: REM-5A05
Phone Number: (571)270-5232
Email: Kristin Bianchi@uspto.gov

Request Detail

Attachment: 569873, Claims, Page Range4 pages.pdf

Case/Application number: 10/569873 PALM

Priority App. Filing Date: 08/29/2003 Format for Search Results: EMAIL

Meaning of unusual acronyms or initialisms:

Identify the novelty:

Additional Comments:

Please search the compounds of claim 1. Please note the proviso on page 3 of the claims (line 9) which states that "there exists at least one of R3 that is halogen or trihalomethyl," Thank you!

Request Date: Tuesday, September 13, 2011 11:12 AM

INVENTOR SEARCH

=> fil capl; d que nos 182 FILE 'CAPLUS' ENTERED AT 14:17:02 ON 14 SEP 2011 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2011 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 14 Sep 2011 VOL 155 ISS 12

FILE LAST UPDATED: 13 Sep 2011 (20110913/ED)

REVISED CLASS FIELDS (/NCL) LAST RELOADED: Jun 2011

USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Jun 2011

CAplus now includes complete International Patent Classification (IPC) reclassification data for the second quarter of 2011.

CAS Information Use Policies apply and are available at:

http://www.cas.org/legal/infopolicy.html

This file contains CAS Registry Numbers for easy and accurate substance identification.

'OBI' IS DEFAULT SEARCH FIELD FOR 'CAPLUS' FILE

L10		STR
L11		STR
L13	5217	SEA FILE=REGISTRY SSS FUL L10 AND L11
L18	165	SEA FILE=CAPLUS SPE=ON ABB=ON L13
L47	4779	SEA FILE=CAPLUS SPE=ON ABB=ON CHENG W?/AU, AUTH
L48	25	SEA FILE=CAPLUS SPE=ON ABB=ON CO E?/AU, AUTH
L49	28337	SEA FILE=CAPLUS SPE=ON ABB=ON KIM M?/AU, AUTH
L50	2799	SEA FILE=CAPLUS SPE=ON ABB=ON KLEIN R?/AU, AUTH
L51	282	SEA FILE=CAPLUS SPE=ON ABB=ON LEW A?/AU OR LEW TSUHAKO A?/AU
		OR TSUHAKO A?/AU, AUTH
L52	204	SEA FILE=CAPLUS SPE=ON ABB=ON NUSS J?/AU, AUTH
L53	16047	SEA FILE=CAPLUS SPE=ON ABB=ON XU W?/AU, AUTH
L54	5	SEA FILE=CAPLUS SPE=ON ABB=ON BAJJALIEH W?/AU, AUTH
L56	11	SEA FILE=CAPLUS SPE=ON ABB=ON L47 AND (L48 OR L49 OR L50 OR
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L57	8	SEA FILE=CAPLUS SPE=ON ABB=ON L48 AND (L49 OR L50 OR L50 OR
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L60	5	L53 OR L54) SEA FILE=CAPLUS SPE=ON ABB=ON L50 AND (L51 OR L52 OR L53 OR L54)
L61	10	SEA FILE=CAPLUS SPE=ON ABB=ON L51 AND (L52 OR L53 OR L54)
L62	14	SEA FILE=CAPLUS SPE=ON ABB=ON L52 AND (L53 OR L54)
L63	2	SEA FILE=CAPLUS SPE=ON ABB=ON L53 AND L54
L65	4	SEA FILE=CAPLUS SPE=ON ABB=ON L56 AND (L57 OR L58 OR L59 OR
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L66	7	SEA FILE=CAPLUS SPE=ON ABB=ON L57 AND (L58 OR L59 OR L60 OR
		L61 OR L62 OR L63)
L67	13	SEA FILE=CAPLUS SPE=ON ABB=ON L58 AND (L59 OR L60 OR L61 OR
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L68	5	SEA FILE=CAPLUS SPE=ON ABB=ON L59 AND (L60 OR L61 OR L62 OR
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L76	5	SEA FILE=CAPLUS SPE=ON ABB=ON L68 AND (L69 OR L70 OR L71)
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_ , ,		L77 OR L78)
L81	1	SEA FILE=CAPLUS SPE=ON ABB=ON (L47 OR L48 OR L49 OR L50 OR
- 	_	L51 OR L52 OR L53 OR L54) AND L18
L82	7	SEA FILE=CAPLUS SPE=ON ABB=ON (L79 OR L81)
	,	

=> d ibib abs hitstr 182 1-7

L82 ANSWER 1 OF 7 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2006:655708 CAPLUS Full-text

DOCUMENT NUMBER: 145:124611 TITLE: Preparation of

[1H-pyrazolo[3,4-d]pyrimidin-4-yl]piperidine or -piperazine compounds as serine-threonine kinase modulators (p70S6K, Akt-1 and Akt-2) for the treatment

of immunological, inflammatory and proliferative

diseases

Rice, Ken; Co, Erick Wang; Kim, Moon Hwan; Bannen, INVENTOR(S):

Lynn Canne; Bussenius, Joerg; Le, Donna; Tsuhako, Amy

Lew; Nuss, John; Wang, Yong; Xu, Wei; Klein,

Rhett Ronald

Exelixis, Inc., USA PATENT ASSIGNEE(S): PCT Int. Appl., 126 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

					KIN					ICAT	DATE						
WO 2006071819																	
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		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KM,	KN,	KP,	KR,
		KΖ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,
		MZ,	NA,	NG,	NI,	NO,	NΖ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,
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ΑU	2005	3220	85		A1		2006	0706		AU 2	005-		20051227				
CA	2590	961			A1		2006	0706	1	CA 2	005-		20051227				
EΡ	1848	719			A1		2007	1031		EP 2	005-	8554	90		2	0051	227
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	2008				_						2007-				_	0051	
US	2008									US 2	2007-	7222	91		2	0071	010
US	7994	172			В2		2011	0809									
RIT	Y APP	LN.	INFO	.:							004-					0041	228
										WO 2	005-	US469	938	1	<i>N</i> 2	0051	227

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): CASREACT 145:124611; MARPAT 145:124611 GI

AB The title compds. I [R1 = H, halo, CN, aryl, etc.; R2 = H, NH2, SH, OH or alkyl; R3-R6 = H, oxo, alkyl, alkoxy, etc.; L = alkylene, alkenylene, C(O), etc.; Q1 = N, CR13 (wherein R13 = H or C(O)NR12(CH2)nNR10R11); Q2 = a bond, CR14, O or N (R14 = H, OH, alkyl, etc.); n = 1-4; W = alkyl, NR10R11, aryl, cycloalkyl, etc.; or V, Q2, L and W together form aryl ring, heteroaryl ring, cycloalkyl ring, etc.; R10, R11, R12 = H or alkyl which is optionally substituted with aryl or heteroaryl; with provisos], useful for inhibition of kinases, more specifically p70S6 kinases, and more preferably p70S6, Akt-1

and Akt-2 kinases, were prepared E.g., a multi-step synthesis of II, starting from N-Boc-4-(4-chlorobenzoyl)piperidine and 2-(diethylamino)ethylamine, was given. Compds. I were tested against p70S6K, Akt-1 and Akt-2 (IC50 values were given for representative compds. I). The invention provides compds. for modulating protein kinase enzymic activity for modulating cellular activities such as proliferation, differentiation, programmed cell death, migration, chemoinvasion and metabolism Compds. I inhibit, regulate and/or modulate kinase receptor signal transduction pathways related to the changes in cellular activities as mentioned above, and the invention includes compns. which contain these compds., and methods of using them to treat kinase-dependent diseases and conditions.

OS.CITING REF COUNT: 7 THERE ARE 7 CAPLUS RECORDS THAT CITE THIS RECORD

(10 CITINGS)

REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L82 ANSWER 2 OF 7 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2005:1314205 CAPLUS Full-text

DOCUMENT NUMBER: 144:51610

TITLE: Preparation and structure activity of

pyrazolo-pyrimidine derivatives as antitumor agents

and kinase modulators

INVENTOR(S): Anand, Neel K.; Blazey, Charles M.; Bowles, Owen

Joseph; Bussenius, Joerg; Canne Bannen, Lynne; Chan,

Diva Sze-Ming; Chen, Baili; Co, Erick Wang;

Costanzo, Simona; Defina, Steven Charles; Dubenko,

Larisa; Franzini, Maurizio; Huang, Ping;

Jammalamadaka, Vasu; Khoury, Richard George; Kim, Moon Hwan; Klein, Rhett Ronald; Le, Donna Tra; Mac, Morrison B.; Nuss, John M.; Parks, Jason Jevious; Rice, Kenneth D.; Tsang, Tsze H.; Tsuhako, Amy Lew;

Wang, Yong; Xu, ₩@i

PATENT ASSIGNEE(S): Exelixis, Inc., USA SOURCE: PCT Int. Appl., 211 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	NO.			KIN	D	DATE			APPLICATION NO.						DATE			
WO 2005		09	A2 20051215 A3 20060427					WO 2005-US13860						20050422				
\overline{W} :	ΑE,	AG,	AL,	AM,	ΑT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,		
	CN,	СО,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,		
	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	\mathtt{KP} ,	KR,	KZ,		
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	ZM,	ZW																
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RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML,
             MR, NE, SN, TD, TG
     AU 2005249380
                          A1
                                 20051215
                                             AU 2005-249380
                                                                     20050422
     CA 2563699
                          A1
                                 20051215
                                             CA 2005-2563699
                                                                     20050422
                                 20070214
                                             EP 2005-804792
     EP 1750727
                          A2
                                                                     20050422
         R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
             IS, IT, LI, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, AL, BA,
             HR, LV, MK, YU
                          Τ
                                 20071129
                                             JP 2007-509678
                                                                     20050422
     JP 2007534687
     US 20080076774
                          A1
                                 20080327
                                             US 2007-568173
                                                                     20070726
PRIORITY APPLN. INFO.:
                                             US 2004-564908P
                                                                  Ρ
                                                                     20040423
                                             WO 2005-US13860
                                                                     20050422
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): CASREACT 144:51610; MARPAT 144:51610 GI

OS.CITING REF COUNT:

16

Pyrazolo-pyrimidine derivs. I, wherein X1 is N, CR2. X2 is N, CR3; X3 is N, AB CR4, but when X2 is N then X3 is CR4; R is H, halogen, tri-halomethyl, substituted nitrogen, substituted sulfur, sulfonyl, sulfonamide, carboxylate, amide, substituted oxygen, acyl, alkyl, aryl, heterocycle, heterocycloalkyl, arylalkyl R1-R13 are independently H, halogen, tri-halomethyl, CN, NO2, substituted nitrogen, substituted sulfur, sulfonyl, sulfonamide, carboxylate, amide, substituted oxygen, acyl, alkyl, aryl, heterocycle, heterocycloalkyl, arylalkyl; Q is (C)nR11R12; n is 0-1 are prepared as kinase modulators. Combination chemotherapy and structure activity of title compds. are reported. The compds. modulate protein kinase enzymic activity to modulate cellular activities such as proliferation, differentiation, programmed cell death, migration and chemoinvasion. Compds. of the invention inhibit, regulate and/or modulate kinases, particularly p70S6 and/or AKT kinases. Methods of using and preparing the compds., and pharmaceutical compns. thereof, to treat kinase-dependent diseases and conditions are also an aspect of the invention. Thus, 3-(azetidin-3-ylidene-methyl)-4-[4-(5-chloro-2-methyl)]methylphenyl)piperazin-1-yl]-1H-pyrazolo[3,4-d]pyrimidine was prepared and tested in vitro as kinase modulator (IC50 > 1000 nM).

THERE ARE 16 CAPLUS RECORDS THAT CITE THIS RECORD (19 CITINGS)

L82 ANSWER 3 OF 7 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2005:395042 CAPLUS Full-text

DOCUMENT NUMBER: 142:447414

TITLE: P70S6 kinase modulators and method of use INVENTOR(S): Cheng, Wei; Co, Erick Wang; Kim, Moon Hwan; Klein, Rhett Ronald; Le Donna, T.; Lew, Amy;

Nuss, John M.; Xu, Wei

Exelixis, Inc., USA PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 165 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

				KIND						APPLICATION NO.									
WC	200	50395	06					WO 2004-US35470											
WC					_			-											
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		•				•		•			EC,			•	•	•			
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		SN,	TD,	ΤG															
JΑ	AU 2004283751				A1 20050506					AU 2004-283751						0041	022		
ΑU	200	42837	51		В2		2011	0519											
CZ	254	1989			A1		2005	0506		CA 2	2004-	2541	989		2	0041	022		
EF	167	8168			A2		2006	0712		EP 2004-796443						20041022			
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		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	PL,	SK, F	ЯŁ	
JE	200	75274																	
US	200	70208	020		A1		2007	0906		US 2	2006-	5766	53		2	0061	116		
	781				В2														
US	201	10021								US 2	2010-	8399	25		2	0100	720		
PRIORIT							-				2003-					0031	024		
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): CASREACT 142:447414; MARPAT 142:447414

GI

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

Peptide derivs. I [E = C(R2)]-substituted pyridine, pyridazine, pyrimidine, AB or 1,3,5-triazine; B = (R1)n; R1, R2 = H, halo, trihalomethyl, CN, NO2, aminoalkyl, carboxyalkyl, (un)substituted alky, alkenyl, alkynyl, aryl, heterocyclyl, heterocyclylalkyl, arylalkyl, etc.; X, Y = CO, O, (un)substituted amine, (un)substituted imine, SO; X and Y can combine to form either C(R3):C(R3), or C.tplbond.C; when X = O, (un)substituted amine, or (un)substituted imine, Y cannot be CH(R3); R3 = (un)substituted Ph, naphthyl, cyclohexyl, dihydronaphthyl, five- to six-membered heteroaryl; Z = 0, S, double bond to an atom of B; A = single bond, NH, (un) substitutedaminoalkyl, aminoaryl, aminoarylalkyl, aminoheterocyclyl, aminoheterocyclylalkyl; J = (un) substituted five- to ten-membered aryl or heteroaryl, etc.; n = 0-5] or pharmaceutically acceptable salts, hydrates, or prodrugs were prepared as p70S6 kinase signal transduction inhibitors and cellular activities modulators for treating kinase-dependent diseases and conditions. Thus, compound II was prepared by coupling of

2-amino-4,6-di-chloro-5-formylpyrimidine with

2-amino-N-(3-trifluoromethylphenyl)acetamide in 43%yield and showed IC50 < 50 nM in p70S6 kinase activity assey.

OS.CITING REF COUNT: 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD

(3 CITINGS)

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L82 ANSWER 4 OF 7 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2005:216619 CAPLUS Full-text

DOCUMENT NUMBER: 142:297864

TITLE: Preparation of aniline derivatives and related

compounds as c-kit modulators

INVENTOR(S): Cheng, Wei; Co, Erick Wang; Kim, Moon Hwan;

Klein, Rhett Ronald; Le Donna, T.; Lew, Amy; Nuss, John M.; Xu, Wei; Bajjalieh, William

PATENT ASSIGNEE(S): Exelixis, Inc., USA SOURCE: PCT Int. Appl., 169 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PA]	ENT 1	. O <i>v</i>			KIND DATE						LICAT								
						A2 20050310 A3 20051006					2004-	20040827								
	W	Z005				AL, AM, AT, AU, AZ, E					BB	. BG.	BR.	BW.	BY.	B7.	CA.	CH.		
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				TD,																
	AU	2004	2686	21		A1 20050310						AU 2004-268621						20040827		
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	CA	25369	954			A1		2005	0310		CA	2004-	2536	954		2	0040	827		
	ΕP	1663	204			A2		2006	0607		EP	2004-	7824	73		2	0040	827		
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	JΡ	2007	5041	60		Τ		2007	0301		JP 2006-524905						20040827			
	US	2008	0096	892		A1		2008	0424		US 2007-569873						20070904			
PRIOR	ΙΤΊ	APP:	LN.	INFO	.:						US 2003-499224P						P 20030829			
											WO	2004-	US28	001		W 2	0040	827		

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): CASREACT 142:297864; MARPAT 142:297864 GI

Compds. I [wherein ring A is a five- to fourteen-membered heteroaryl; R1, R2 and R3 are H, halo, trihalomethyl, cyano, nitro, etc.; L1 is a single bond, (un) substituted alkylene, O, CH2O, etc.; ring B is five- to ten-membered aryl or heterocyclyl; ring C is five- to ten-membered (hetero)aryl; L2 is alkylene, alkylidene, alkylidyne, etc.; with some limitations and exclusions, and pharmaceutically acceptable salts, hydrates or prodrugs thereof], as exemplified by carbonyl compds. of anilines, were prepared as c-Kit kinase modulators. For example, 3-aminophenoxyacetic acid, which was obtained from the corresponding nitro compound in 76% yield via catalytic hydrogenation, was treated with HC(OEt)3 and NaN3 in AcOH followed by NaNO2/HCl to give a tetrazole in 61% yield. This acid was coupled with 5-amino-2-chlorobenzotrifluoride in the presence of HATU to afford acetamide

II in 46% yield, which showed inhibition against c-Kit kinase with a IC50 of <50 nM. Therefore, I and pharmaceutical compns. thereof are useful for modulating c-Kit kinase activity and for treating diseases or disorders associated with uncontrolled, abnormal, and/or unwanted cellular activities.

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ΙT
     332176-74-69
                    483337-32-2P
                                    483337-34-4P
     483337-36-6P
                    483337-37-79
                                    483337-38-82
                                    483337-41-3P
     483337-39-97
                    483337-40-29
                    505052-18-6P
     483978-03-6P
                                    506433-09-69
     552825-29-39
                    847606-67-19
                                    847606-71-7P
     847606-73-92
                    847606-74-0P
                                    847606-76-2P
     847606-77-3P
                   847606-81-92
                                    847606-84-2P
     847606-87-5P
                    847606-90-02
                                    847606-93-3P
     847606-95-5P
                    847607-13-0P
                                    847607-14-12
     847607-15-22
                    847607-17-49
                                    847607-18-5P
     847607-19-6P
                    847607-20-99
                                    847607-22-1P
     847607-25-4P
                    847607-26-59
                                    847607-27-6P
     847607-28-72
                    847607-29-89
                                    847607-37-8P
     847607-38-92
                    847607-47-09
                                    847607-48-19
     847607-51-6P
                    847607-58-3P
                                    847607-61-8P
     847607-68-5P
                    847607-69-6P
                                    847607-71-0P
     847607-74-3P
                                    847607-77-6P
                    847607-76-5P
     847607-78-72
                    847607-79-8P
                                    847607-80-19
     847607-81-2P
                    847607-82-32
                                    847607-83-42
     847607-84-59
                    847607-86-72
                                    847607-87-82
     847607-88-9P
                    847607-89-0P
                                    847607-92-52
     847607-93-69
                    847607-94-72
                                    847607-95-82
     847607~96~99
                    847607-97-02
                                    847607-98-12
     847607-99-22
                    847608-00-89
                                    847608-01-9P
     847608-02-0P
                    847608-03-19
                                    847608-04-29
     847608-05-3P
                    847608-06-4P
                                    847608-07-5P
     847608-08-6P
                    847608-09-7P
                                    847608-10-0P
     847608-12-28
                    847608-13-3P
                                    847608-14-42
     847608-15-5P
                    847608-16-62
                                    847608-17-72
     847608-18-89
                    847608-19-92
                                    847608-20-2P
     847608-21-3P
                    847608-23-59
                                    847608-24-69
     847608-25-7P
                    847608-26-8P
                                    847608-27-92
     847608-29-19
                    847608-30-4P
                                    847608-31-5P
     847608-32-6P
                    847608-33-79
                                    847608-35-99
     847608-37-1P
                    847608-39-3P
                                    847608-42-89
     847608-44-0P
                    847608-46-2P
                                    847608-48-4P
                                    847608-53-1P
     847608-50-8P
                    847608-51-9P
     847608-55-3P
                    847608-58-6P
                                    847608-59-79
     847608-60-0P
                    847608-62-2P
                                    847608-63-3P
     847608-65-5P
                    847608-66-62
                                    847608-67-7P
     847608-68-8P
                                    847608-70-2P
                    847608-69-9P
     847608-71-3P
                    847608-74-6P
                                    847608-75-7P
     847608-77-99
                    847608-79-1P
                                    847608-80-49
     847608-81-5P
                    847608-82-6P
                                    847608-83-7P
     847608-84-8P
                    847608-85-9P
                                    847608-86-0P
     847608-87-1P
                    847608-89-3P
                                    847608-90-6P
     847608-91-72
                    847608-98-4P
                                    847609-00-1P
     847609-04-5P
                    847609-06-7P
                                    847609-08-9F
     847609-10-3P
                    847609-12-5P
                                    847609-14-72
     847609-18-19
                    847609-20-5P
                                    847609-28-3P
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847609-30-79 847609-32-99 847609-50-1P 847609-52-3P 847609-54-5P 847609-56-79 847609-57-8P 847609-58-9P 847609-59-0P 847609-60-3P 847609-63-6P 847609-65-8P 847609-67-09 847609-73-8P 847609-75-0P 847609-81-89 847609-79-4P 847609-86-3P 847609-93-29

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(modulator; preparation of anilines and related compds. as C-kit
modulators)

RN 332176-74-6 CAPLUS

CN Acetamide, N-(5-chloro-2-methoxyphenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 483337-32-2 CAPLUS

CN Acetamide, N-(2-chlorophenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c} N \\ N \\ \end{array} \begin{array}{c} N \\ \end{array} \begin{array}{c} O \\ \end{array} \begin{array}{c} C \\ H \\ \end{array} \begin{array}{c} O \\ \end{array} \begin{array}$$

RN 483337-34-4 CAPLUS

CN Acetamide, N-(2,3-dichlorophenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ &$$

RN 483337-36-6 CAPLUS

CN Acetamide, N-(3-chloro-2-methylphenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 483337-37-7 CAPLUS

CN Acetamide, N-(4-bromophenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 483337-38-8 CAPLUS

CN Acetamide, N-(2-fluorophenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 483337-39-9 CAPLUS

CN Acetamide, N-(4-fluorophenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 483337-40-2 CAPLUS

CN Acetamide, 2-[3-(1H-tetrazol-1-yl)phenoxy]-N-[2-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 483337-41-3 CAPLUS

CN Acetamide, 2-[3-(1H-tetrazol-1-y1)phenoxy]-N-[3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 483978-03-6 CAPLUS

CN Acetamide, N-(3-fluorophenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 505052-18-6 CAPLUS

CN Acetamide, N-[2-chloro-5-(trifluoromethyl)phenyl]-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 506433-09-6 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$N = 0 - CH_2 - C - NH - C1$$

RN 552825-29-3 CAPLUS

CN Acetamide, N-(4-chlorophenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847606-67-1 CAPLUS

CN Acetamide,

N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(1H-1,2,3-triazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 847606-71-7 CAPLUS

CN Hydrazinecarboxamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(1H-tetrazol-1-yl)phenyl]- (CA INDEX NAME)

RN 847606-73-9 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[[3-(1H-tetrazol-1-yl)phenyl]amino]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 847606-74-0 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(5-methyl-1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} N & & \\ N & & \\ N & & \\ \end{array}$$

RN 847606-76-2 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[[3-(1H-tetrazol-1-yl)phenyl]thio]- (CA INDEX NAME)

$$N = N = CH_2 - 0 = CH_2 - 0$$

$$CF_3 = CH_2 - 0$$

RN 847606-77-3 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(5-pyrimidinyl)phenoxy]- (CA INDEX NAME)

RN 847606-81-9 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(1H-tetrazol-1-yl)phenyl]methyl]- (CA INDEX NAME)

RN 847606-84-2 CAPLUS

CN Propanamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847606-87-5 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(6-quinoxalinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847606-90-0 CAPLUS

CN Benzamide, N-[[[4-chloro-3-(trifluoromethyl)phenyl]amino]carbonyl]-3-(1H-tetrazol-1-yl)- (CA INDEX NAME)

RN 847606-93-3 CAPLUS

CN Carbamic acid, [3-(trifluoromethyl)phenyl]-, [3-(1H-tetrazol-1-yl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847606-95-5 CAPLUS

CN Acetamide, N-(4-chloro-3-methylphenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-13-0 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(2H-tetrazol-5-yl)phenoxy]- (CA INDEX NAME)

RN 847607-14-1 CAPLUS

CN Acetamide, N-(4-chloro-2-fluorophenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-15-2 CAPLUS

CN Acetamide, N-(4-bromo-3-methylphenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-17-4 CAPLUS

CN Acetamide, N-[4-fluoro-3-(trifluoromethyl)phenyl]-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-18-5 CAPLUS

CN Acetamide, N-[4-bromo-3-(trifluoromethyl)phenyl]-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} \mathbb{N} & \mathbb{N} &$$

RN 847607-19-6 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-20-9 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[2-methyl-5-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} & \text{Me} & \text{O} \\ & \text{O} & \text{CH}_2 - \text{C} \\ & \text{NH} \end{array}$$

RN 847607-22-1 CAPLUS

CN Acetamide, N-[4-chloro-2-(trifluoromethyl)phenyl]-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 847607-25-4 CAPLUS

CN Acetamide, N-[4-fluoro-3-(trifluoromethyl)phenyl]-2-[4-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-26-5 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(2-methyl-2H-tetrazol-5-yl)phenoxy]- (CA INDEX NAME)

Me N
$$\sim$$
 N \sim CH₂ \sim NH \sim C1

RN 847607-27-6 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[2,4-dichloro-5-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 847607-28-7 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[2-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-29-8 CAPLUS

CN 1H-1,2,3-Triazole-4-carboxylic acid,

1-[3-[2-[[4-chloro-3-(trifluoromethyl)phenyl]amino]-2-oxoethoxy]phenyl]-, methyl ester (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 847607-37-8 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(4-pyridinyl)phenoxy]- (CA INDEX NAME)

RN 847607-38-9 CAPLUS

CN Propanamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[[3-(1H-tetrazol-1-yl)phenyl]amino]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 847607-47-0 CAPLUS

CN Benzamide, 5-chloro-2-[[2-[3-(1H-tetrazol-1-yl)phenoxy]acetyl]amino]-

(CA

INDEX NAME)

RN 847607-48-1 CAPLUS

CN Acetamide, N-(5-chloro-2,4-dimethoxyphenyl)-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-51-6 CAPLUS

CN Acetamide, N-[2-methoxy-5-(trifluoromethyl)phenyl]-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-58-3 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(5-pyrimidinyl)phenoxy]- (CA INDEX NAME)

RN 847607-61-8 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(3-pyridinyl)phenoxy]- (CA INDEX NAME)

RN 847607-68-5 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(3,5-dimethyl-4-isoxazolyl)phenoxy]- (CA INDEX NAME)

RN 847607-69-6 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(7-quinolinyl)phenoxy]- (CA INDEX NAME)

RN 847607-71-0 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(4-dibenzofuranyl)phenoxy]- (CA INDEX NAME)

RN 847607-74-3 CAPLUS

CN Acetamide, N-[4-fluoro-3-(trifluoromethyl)phenyl]-2-[[3-(1H-tetrazol-1-yl)phenyl]amino]- (CA INDEX NAME)

$$N = NH - CH_2 - C - NH - CH_3 - CH_3$$

RN 847607-76-5 CAPLUS

CN Hydrazinecarboxamide, N-[2-fluoro-5-(trifluoromethyl)phenyl]-2-[3-(1H-tetrazol-1-yl)phenyl]- (CA INDEX NAME)

RN 847607-77-6 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(3-pyridinyl)phenoxy]- (CA INDEX NAME)

RN 847607-78-7 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(5-pyrimidinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847607-79-8 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethy1)pheny1]-N'-[[4-(5-pyrimidiny1)pheny1]methy1]- (CA INDEX NAME)

RN 847607-80-1 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & \circ & \\ \text{CH}_2 - \text{NH} - \overset{\circ}{\text{C}} - \text{NH} \\ & & \text{CI} \end{array}$$

RN 847607-81-2 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(1H-tetrazol-1-yl)phenyl]methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \mathbb{N} & \mathbb{N} & \mathbb{C} \\ \mathbb{N} & \mathbb{C} \\ \mathbb{C} \\ \mathbb{C} & \mathbb{C} \\ \mathbb{C} \\ \mathbb{C} & \mathbb{C} \\ \mathbb{C} \\ \mathbb{C} & \mathbb{C} \\ \mathbb{C} \\ \mathbb{C} \\ \mathbb{C} & \mathbb{C} \\ \mathbb{C} \\$$

RN 847607-82-3 CAPLUS

CN Acetamide, N-[4-fluoro-3-(trifluoromethyl)phenyl]-2-[4-(5-pyrimidinyl)phenoxy]- (CA INDEX NAME)

$$0 - CH_2 - C - NH - CF_3$$

RN 847607-83-4 CAPLUS

CN Acetamide,

2-[[4-chloro-3-(trifluoromethyl)phenyl]amino]-N-[3-(1H-tetrazol-

1-yl)phenyl]- (CA INDEX NAME)

RN 847607-84-5 CAPLUS

CN Acetamide, 2-[4-chloro-3-(trifluoromethyl)phenoxy]-N-[3-(1H-tetrazol-1-yl)phenyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 847607-86-7 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-methyl-4-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-87-8 CAPLUS

CN Acetamide,

N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(1H-1,2,3-triazol-1-yl)phenoxy]- (CA INDEX NAME)

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-fluoro-4-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-89-0 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[2-fluoro-4-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847607-92-5 CAPLUS

CN Acetamide, N-[4-fluoro-3-(trifluoromethyl)phenyl]-2-[4-(3-pyridinyl)phenoxy]- (CA INDEX NAME)

RN 847607-93-6 CAPLUS

CN Acetamide, 2-[4-(2,4-dimethoxy-5-pyrimidinyl)phenoxy]-N-[4-fluoro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 847607-94-7 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(2,4-dimethoxy-5-pyrimidinyl)phenoxy]- (CA INDEX NAME)

RN 847607-95-8 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(4-pyridinyl)phenoxy]- (CA INDEX NAME)

RN 847607-96-9 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[[3-methoxy-4-(1H-tetrazol-1-yl)phenyl]amino]- (CA INDEX NAME)

RN 847607-97-0 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[[4-methoxy-3-(1H-tetrazol-1-yl)phenyl]amino]- (CA INDEX NAME)

$$\begin{array}{c} \text{NM=O} \\ \text{NM=CH2-C-NH-CH3} \\ \text{CF3} \end{array}$$

RN 847607-98-1 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[[4-(1H-tetrazol-1-yl)phenyl]amino]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 847607-99-2 CAPLUS

CN Hydrazinecarboxamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[2,3,5,6-tetrafluoro-4-(5-pyrimidinyl)phenyl]- (CA INDEX NAME)

RN 847608-00-8 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(1H-tetrazol-1-yl)phenyl]methyl]- (CA INDEX NAME)

$$\begin{array}{c} N \\ N \\ \end{array}$$

RN 847608-01-9 CAPLUS

CN Hydrazinecarboxamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(5-pyrimidinyl)phenyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &$$

RN 847608-02-0 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-03-1 CAPLUS

CN Propanamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-methyl-2-[3-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847608-04-2 CAPLUS

CN Propanamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

RN 847608-05-3 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(2,4-dimethoxy-5-pyrimidinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-06-4 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(2-methoxy-5-pyrimidinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-07-5 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(6-methoxy-3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-08-6 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(2-methoxy-5-pyrimidinyl)phenyl]methyl]- (CA INDEX NAME)

$$\begin{array}{c} \text{MeO} \\ \text{N} \end{array}$$

RN 847608-09-7 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(6-methoxy-3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-10-0 CAPLUS

CN 1H-Indole-1-carboxylic acid, 2-[4-[2-[[4-chloro-3-(trifluoromethyl)phenyl]amino]-2-oxoethoxy]phenyl]-, 1,1-dimethylethyl ester (CA INDEX NAME)

RN 847608-12-2 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[[3-(2H-tetrazol-5-yl)phenyl]amino]- (CA INDEX NAME)

RN 847608-13-3 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[2,6-difluoro-4-(1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 847608-14-4 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-15-5 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(5-pyrimidinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-16-6 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(4-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-17-7 CAPLUS

CN Hydrazinecarboxamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(1H-tetrazol-1-yl)phenyl]- (CA INDEX NAME)

RN 847608-18-8 CAPLUS

CN Hydrazinecarboxamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(3-pyridinyl)phenyl]- (CA INDEX NAME)

RN 847608-19-9 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [4-(3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-20-2 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [4-(4-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-21-3 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [4-(5-pyrimidinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-23-5 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(4-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ &$$

RN 847608-24-6 CAPLUS

CN Hydrazinecarboxamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(3-pyridinyl)phenyl]- (CA INDEX NAME)

RN 847608-25-7 CAPLUS

CN Hydrazinecarboxamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(5-pyrimidinyl)phenyl]- (CA INDEX NAME)

RN 847608-26-8 CAPLUS

CN Urea, N-(5-chloro-2,4-dimethoxyphenyl)-N'-[[4-(5-pyrimidinyl)phenyl]methyl]- (CA INDEX NAME)

$$\begin{array}{c} \text{OMeO} \\ \text{NH} \end{array}$$

RN 847608-27-9 CAPLUS

CN Urea,

N-(5-chloro-2,4-dimethoxyphenyl)-N'-[[4-(3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-29-1 CAPLUS

CN Carbamic acid, (5-chloro-2,4-dimethoxyphenyl)-, [4-(5-pyrimidinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-30-4 CAPLUS

CN Carbamic acid, (5-chloro-2,4-dimethoxyphenyl)-, [4-(3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

$$\text{CH}_2\text{-}\text{O}\text{-}\text{NH}\text{-}\text{Cl}$$

RN 847608-31-5 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, 1-[4-(3-pyridinyl)phenyl]ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Me} & \text{O} \\ \text{CH-O-C-NH} \\ \text{CF}_3 \end{array}$$

RN 847608-32-6 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, 1-[4-(5-pyrimidinyl)phenyl]ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Me} & \text{O} \\ \text{CH} & \text{O} & \text{C} \\ \text{NH} & \text{CI} \end{array}$$

RN 847608-33-7 CAPLUS

CN Urea,

N-(5-chloro-2,4-dimethoxyphenyl)-N'-[[3-(3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & \\ & & \\$$

RN 847608-35-9 CAPLUS

CN Urea, N-(5-chloro-2,4-dimethoxyphenyl)-N'-[[3-(5-pyrimidinyl)phenyl]methyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ \end{array}$$

RN 847608-37-1 CAPLUS

CN Carbamic acid, (5-chloro-2,4-dimethoxyphenyl)-,
[3-(3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-39-3 CAPLUS

CN Carbamic acid, (5-chloro-2,4-dimethoxyphenyl)-, [3-(5-pyrimidinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-42-8 CAPLUS

CN Urea, N-[4-fluoro-3-(trifluoromethyl)phenyl]-N'-[[4-(3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-44-0 CAPLUS

CN Urea, N-[[3-(6-amino-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 847608-46-2 CAPLUS

CN Urea, N-[[4-(6-amino-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 847608-48-4 CAPLUS

CN Urea, N-[[3-(2-amino-5-pyrimidinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

RN 847608-50-8 CAPLUS

CN Urea, N-[[4-(2-amino-5-pyrimidinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

$$\begin{array}{c} \text{CF}_3 \\ \text{H}_2\text{N} \\ \text{N} \end{array} \begin{array}{c} \text{CH}_2 - \text{NH} \\ \text{C} - \text{NH} \end{array} \begin{array}{c} \text{CF}_3 \\ \text{C}_1 \\ \text{C}_1 \\ \text{C}_2 \\ \text{C}_3 \\ \text{C}_1 \\ \text{C}_1 \\ \text{C}_2 \\ \text{C}_3 \\ \text{C}_1 \\ \text{C}_2 \\ \text{C}_3 \\ \text{C}_1 \\ \text{C}_2 \\ \text{C}_3 \\ \text{C}_1 \\ \text{C}_4 \\ \text{C}_4 \\ \text{C}_5 \\ \text{C}_1 \\ \text{C}_2 \\ \text{C}_3 \\ \text{C}_4 \\ \text{C}_5 \\ \text{C$$

RN 847608-51-9 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[1-[4-(3-pyridinyl)phenyl]ethyl]- (CA INDEX NAME)

$$\begin{array}{c} \text{Me} & \text{O} \\ \text{CH-NH-C-NH} \\ \text{CF}_3 \end{array}$$

RN 847608-53-1 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[1-[4-(5-pyrimidinyl)phenyl]ethyl]- (CA INDEX NAME)

$$\begin{array}{c} \text{Me} \\ \text{CH-NH-C-NH} \\ \end{array}$$

RN 847608-55-3 CAPLUS

CN Acetamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(1H-indol-2-yl)phenoxy]- (CA INDEX NAME)

RN 847608-58-6 CAPLUS

CN Hydrazinecarboxamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(4-pyridinyl)phenyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & \circ & \circ \\ & \text{NH-NH-C-NH} \\ & & \text{CF}_3 \end{array}$$

RN 847608-59-7 CAPLUS

CN Hydrazinecarboxamide, N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(4-pyridinyl)phenyl]- (CA INDEX NAME)

RN 847608-60-0 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(4-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-62-2 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(6-quinoxalinyl)phenyl]methyl]- (CA INDEX NAME)

$$C1$$
 NH
 C
 NH
 C
 NH
 CH_2
 N

RN 847608-63-3 CAPLUS

CN Urea, N-[[3-(2-amino-5-methyl-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 847608-65-5 CAPLUS

CN Carbamic acid, (3-chloro-4-methoxyphenyl)-,

[3-(1H-tetrazol-1-y1)phenyl]methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} N & & \\ N & & \\ N & & \\ \end{array}$$

RN 847608-66-6 CAPLUS

CN Urea,

N-(3-chloro-4-methoxyphenyl)-N'-[[3-(1H-tetrazol-1-yl)phenyl]methyl]- (CA INDEX NAME)

$$\begin{array}{c|c} N & & \\ N & & \\ \end{array}$$
 CH₂-NH-C-NH-C-NH-OMe

RN 847608-67-7 CAPLUS

CN Acetamide,

N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[4-(2,5-dihydro-5-oxo-1H-tetrazol-1-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 847608-68-8 CAPLUS

CN Urea, N-[[3-(2-amino-5-chloro-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

$$\begin{array}{c} \text{NH}_2 \\ \text{N} \\ \text{C}_1 \end{array} \quad \begin{array}{c} \text{CF}_3 \\ \text{C}_1 \\ \text{C}_1 \end{array} \quad \begin{array}{c} \text{CF}_3 \\ \text{C}_1 \\ \text{C}_1 \\ \text{C}_1 \end{array} \quad \begin{array}{c} \text{CF}_3 \\ \text{C}_1 \\ \text{C}_2 \\ \text{C}_2 \\ \text{C}_2 \\ \text{C}_2 \\ \text{C}_2 \\ \text{C}_2 \\ \text{C}_3 \\ \text{C}_4 \\ \text{C}_2 \\ \text{C}_4 \\ \text{C}_2 \\ \text{C}_4 \\ \text{C}_4 \\ \text{C}_5 \\ \text{C}_5 \\ \text{C}_5 \\ \text{C}_6 \\ \text{C}_6 \\ \text{C}_6 \\ \text{C}_7 \\$$

RN 847608-69-9 CAPLUS

CN Urea, N-[[4-(2-amino-5-chloro-3-pyridiny1)pheny1]methy1]-N'-[4-chloro-3-(trifluoromethy1)pheny1]- (CA INDEX NAME)

RN 847608-70-2 CAPLUS

CN Urea, N-[[3-(6-chloro-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 847608-71-3 CAPLUS

CN Urea, N-[[4-(6-chloro-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 847608-74-6 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(6-fluoro-3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-75-7 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(2-methoxy-3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-77-9 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(6-fluoro-3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-79-1 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(2-methoxy-3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-80-4 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(6-methyl-3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-81-5 CAPLUS

CN Urea, N-[[4-(2-amino-5-fluoro-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 847608-82-6 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(6-methyl-3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-83-7 CAPLUS

CN Urea, N-[[4-(2-amino-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

$$CF_3$$
 C1

RN 847608-84-8 CAPLUS

CN Urea, N-[[3-(2-amino-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 847608-85-9 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(6-methyl-3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-86-0 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(2-amino-5-fluoro-3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-87-1 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(2-amino-3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847608-89-3 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-[6-(hydroxymethyl)-3-pyridinyl]phenyl]methyl]- (CA INDEX NAME)

RN 847608-90-6 CAPLUS

CN Urea, N-[[3-(6-acetyl-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 847608-91-7 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(6-cyano-3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847608-98-4 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(1H-pyrrolo[2,3-d]pyrimidin-4-yl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847609-00-1 CAPLUS

CN Urea, N-[[3-(2-amino-5-fluoro-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 847609-04-5 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(1H-benzimidazol-2-yl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847609-06-7 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(6-amino-2-methyl-3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Me} \\ \text{N} \\ \text{H2N} \end{array} \begin{array}{c} \text{CH}_2 \\ \text{O} \\ \text{CH}_2 \end{array} \begin{array}{c} \text{CF}_3 \\ \text{C1} \\ \text{CH}_2 \end{array}$$

RN 847609-08-9 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-[5-(methylthio)-3-pyridinyl]phenyl]methyl]- (CA INDEX NAME)

RN 847609-10-3 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [4-(6-methyl-3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{CF}_3 \\ \text{CH}_2 - \text{O} \\ \text{C} - \text{NH} \end{array}$$

RN 847609-12-5 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [4-(2-amino-5-fluoro-3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847609-14-7 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [4-(2-amino-3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847609-18-1 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [4-(1H-pyrrolo[2,3-d]pyrimidin-4-yl)phenyl]methyl ester (9CI) (CA INDEX NAME)

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RN 847609-20-5 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [4-(6-amino-2-methyl-3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{Me} \\ \text{H2N} \end{array}$$

RN 847609-28-3 CAPLUS

CN Carbamic acid, (5-chloro-2-methoxyphenyl)-, [3-(3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847609-30-7 CAPLUS
CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-,
[4-(1H-tetrazol-1-yl)phenyl]methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

RN 847609-32-9 CAPLUS

CN Carbamic acid, (5-chloro-2-methoxyphenyl)-, [3-(5-pyrimidinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847609-50-1 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(1H-pyrazol-4-yl)phenyl]methyl]- (CA INDEX NAME)

RN 847609-52-3 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(1H-pyrazol-4-

yl)phenyl]methyl]- (CA INDEX NAME)

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 d_1

RN 847609-54-5 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-[2-(1-piperazinyl)-5-pyrimidinyl]phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847609-56-7 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [4-[2-(1-piperazinyl)-5-pyrimidinyl]phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847609-57-8 CAPLUS

CN Urea, N-[[3-(2-chloro-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

RN 847609-58-9 CAPLUS

CN Urea, N-[[4-(2-chloro-3-pyridinyl)phenyl]methyl]-N'-[4-chloro-3-(trifluoromethyl)phenyl]- (CA INDEX NAME)

$$C1$$
 CH_2 NH CH_2 NH CH_2 CH_3 CH_4 CH_4 CH_5 CH_5 CH_5 CH_5 CH_6 CH_6 CH_6 CH_7 CH_8 CH_8

RN 847609-59-0 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(2-fluoro-3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847609-60-3 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(2-fluoro-3-pyridinyl)phenyl]methyl]- (CA INDEX NAME)

$$\begin{array}{c} CF_3 \\ CH_2-NH \\ \end{array}$$

RN 847609-63-6 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-[5-(methylthio)-2-pyridinyl]phenyl]methyl]- (CA INDEX NAME)

RN 847609-65-8 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(2,6-dimethyl-3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847609-67-0 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(5-methoxy-3-pyridinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847609-73-8 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[3-(4-isoquinolinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847609-75-0 CAPLUS

CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[[4-(4-isoquinolinyl)phenyl]methyl]- (CA INDEX NAME)

RN 847609-79-4 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [3-(1H-pyrazol-4-yl)phenyl]methyl ester (9CI) (CA INDEX NAME)

RN 847609-81-8 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [4-(1H-pyrazol-4-yl)phenyl]methyl ester (9CI) (CA INDEX NAME)

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PAGE 2-A

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RN 847609-86-3 CAPLUS

CN Acetamide,

 $\begin{tabular}{ll} N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-(2,5-dihydro-5-oxo-1H-tetrazol-1-yl)phenoxy]- & (CA INDEX NAME) \end{tabular}$

$$\begin{array}{c|c} & & & \\ & & &$$

RN 847609-93-2 CAPLUS

CN Carbamic acid, [4-chloro-3-(trifluoromethyl)phenyl]-, [4-(4-pyrimidinyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \circ & \\ &$$

IT 847606-70-6P

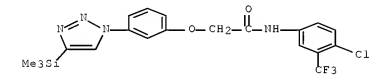
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of anilines and related compds. as C-kit modulators)

RN 847606-70-6 CAPLUS

CN Acetamide,

N-[4-chloro-3-(trifluoromethyl)phenyl]-2-[3-[4-(trimethylsilyl)-1H-1,2,3-triazol-1-yl]phenoxy]- (CA INDEX NAME)



OS.CITING REF COUNT: 7 THERE ARE 7 CAPLUS RECORDS THAT CITE THIS RECORD

(8 CITINGS)

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L82 ANSWER 5 OF 7 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2004:493723 CAPLUS Full-text

DOCUMENT NUMBER: 141:54195

TITLE: Preparation of oxindole derivatives as kinase

modulators

INVENTOR(S): Bannen, Lynne Canne; Brown, S. David; Cheng, Wei;

Co. Erick Wang; Nuss, John M.; Kim, Moon Hwan; Klein, Rhett Ronald; Le, Donna T.; Lew, Amy; Mac, Morrison B.; Parks, Jason Jevious; Wen, Zhaoyang; Xu,

Wei

PATENT ASSIGNEE(S): Exelixis, Inc., USA

SOURCE: PCT Int. Appl., 120 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	ENT	NO.		KIN	D	DATE			APPL	DATE								
						_												
WO 2004050681				A2		2004	0617	,	WO 2	003-1	US36	567		2	0031	114		
WO 2004050681				A3		2004	1104											
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	

CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE,

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GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
             LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ,
             OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM,
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PRIORITY APPLN. INFO.:
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                                                                 Ρ
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                                                                     20030514
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                                                                     20031114
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 141:54195

GΙ

AB The title compds. I [W = N or CR1; R1 = H, halo, trihaloalkyl, CN, NH2, NO2, OR6, N=CNR6R7, N(R6)C(=NR8)NR6R7, SR6, S(0)1-2R6, SO2NR6R7, CO2R6, etc.; L = O, S(0)0-2, or NR3; Q = C or N, when Q = N, then R4 does not exist; R2, R3 = H or R7; R4, R5 = H, OR6, NR6R7, S(0)0-2R6, SO2NR6R7, CO2R6, C(0)NR6R7, N(R6)SO2R6, NC(0)2R6, C(0)R7, CN, NO2, NH2, halo, trihaloalkyl, R7; or R4, R5 when taken together, form a five or six-membered aromatic ring containing 0-2 N; R6, R7 = H, (substituted)(aryl)alkyl, (substituted)heterocyclylalkyl, (substituted)aryl,

(substituted) heterocyclyl, with proviso or R6, R7 = when taken together with a common N to which they are attached, form a five to seven-membered heterocyclic ring containing at least one addnl. heteroatom selected from N, O, S, or P; R8 = H, NO2, CN, OR6, or (substituted) alkyl; X = (substituted) (hetero) aromatic ring; K = O, S, (substituted) amino] were prepared as kinase modulators to treat kinase-dependent diseases and conditions. For example compound II was prepared in a multi-step synthesis starting from 4-methylimidazole. The latter inhibited KDR and EGFR with IC50 < 50 nM.

OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD

(1 CITINGS)

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L82 ANSWER 6 OF 7 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2003:1006921 CAPLUS Full-text

DOCUMENT NUMBER: 140:42210

TITLE: Preparation of 1-sulfonyl-2-piperazinehydroxamic acids

as selective inhibitors of human ADAM-10 for treating cancer, arthritis and diseases related to angiogenesis

INVENTOR(S): Bannen, Lynne Canne; Co, Erick W.; Jammalamadaka,

Vasu; Nuss, John M.; Kim, Moon Hwan; Le Tra,

Donna; Xew, Amy; Mac, Morrison B.; Mamo, Shumeye;

Wen, Zhaoyang; Xu, Wei

PATENT ASSIGNEE(S): Exelixis, Inc., USA

SOURCE: PCT Int. Appl., 94 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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US 7 62 9341	B2	20091208				
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US 79 8966 1	B2	20110802				
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JP 2010265276	A	20101125	JP	2010-142096		20100622
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			JP	2004-513217	A 3	20030611
			MO	2003-US18262	\mathbb{W}	20030611
			US	2005-518110	A 3	20051026

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 140:42210
GI

The present invention provides 1-sulfonyl-2-piperazinehydroxamic acids (shown as I; variables defined below; e.g. II) useful for inhibiting the ADAM-10 protein, with selectivity vs. MMP-1. Inhibition activities of 66 examples of I towards ≤8 metalloproteinases are tabulated. Such compds. are useful in the in vitro study of the role of ADAM-10 (and its inhibition) in biol. processes. The present invention also comprises pharmaceutical compns. comprising ≥1 ADAM-10 inhibitors according to the invention in combination with a pharmaceutically acceptable carrier. Such compns. are useful for the treatment of cancer, arthritis, and diseases related to angiogenesis. Correspondingly, the invention also comprises methods of treating forms of cancer, arthritis, and diseases related to angiogenesis in which ADAM-10 plays a critical role. A method of preparation of sulfonyl halide intermediates is claimed. For example,

[4-(4-fluorophenoxy)-3,5-difluorophenyl]sulfonyl chloride was prepared in 3 steps (105, 98 and 83 % yields) starting from 3,4,5-trifluoronitrobenzene, 4-fluorophenol, and Cs2CO3 in DMF and involving intermediates

- 4-(4-fluorophenoxy)-3,5-difluoronitrobenzene and
- 4-(4-fluorophenoxy)-3,5-difluoroaniline. The prepared
- [4-(4-fluorophenoxy)-3,5-difluorophenyl] sulfonyl chloride was used in a 5-step procedure (65, 78, -, 69 and 62 % yields) to give II involving intermediates
- (R)-1-[[4-(4-fluorophenoxy)-3,5-difluorophenyl]sulfonyl]-4-boc-piperazine-2-carboxylic acid, Me

(R)-1-[[4-(4-fluorophenoxy)-3,5-difluorophenyl]sulfonyl]-4-boc-piperazine- 2-carboxylate, Me (R)-1-[[4-(4-fluorophenoxy)-3,5difluorophenyl]sulfonyl]piperazine-2-carboxylate trifluoroacetate and Me (R)-1-[[4-(4-fluorophenoxy)-3,5-difluorophenyl]sulfonyl]-4-(ethoxycarbonyl)piperazine-2-carboxylate. Although the methods of preparation of I are not claimed, several example prepns. and characterization data for 66 examples of I are included. For I: L1 is -C(0)-, -S(0)2-, or -(CH2)n-; R1 is -H, -OR11, -(CH2)nR11, -C(0)R11, or -NR12R13; R2 is -R21-L2-R22 (R21 is saturated or mono- or poly- unsatd. C5-C14-monoor fused poly-cyclic hydrocarbyl, optionally containing one or two annular heteroatoms per ring and (un)substituted with 1-3 R50 substituents; L2 is -O-, -C(O)-, -CH2-, -NH-, -SO2- or a direct bond; R22 is saturated or monoor poly- unsatd. C5-C14-mono- or fused polycyclic hydrocarbyl, optionally containing one or two annular heteroatoms per ring and (un) substituted with 1-3 R50 substituents); n = 0-3; provided that an O or S is not singly bonded to another O or S in a chain of atoms; addnl. details are given in the claims.

OS.CITING REF COUNT: 7 THERE ARE 7 CAPLUS RECORDS THAT CITE THIS RECORD

(8 CITINGS)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L82 ANSWER 7 OF 7 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2003:491172 CAPLUS Full-text

DOCUMENT NUMBER: 139:69520

TITLE: Preparation of N-sulfonyl amino acid hydroxamide

derivatives as human ADAM-10 inhibitors

INVENTOR(S): Brown, S. David; Canne, Lynne; Co, Erick W.;

Jammalamadaka, Vasu; Khoury, Richard G.; Kim, Moon Hwan; Le, Donna T.; Lew, Amy; Mac, Morrison B.; Mamo, Shumeye; Nuss, John M.; Prisbylla, Michael P.;

Xu, Wei

PATENT ASSIGNEE(S): Exelixis, Inc., USA SOURCE: PCT Int. Appl., 144 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 139:69520

The invention provides amino acid derivs. R5SO2NR4CHR3CONR2OR1 [R1 is H, alkyl, alkanoyl, (un)substituted arylalkyl or arylalkanoyl; R2 is any group given for R1 plus alkoxy; R3 is -Z-Q-J, where Z is (un)substituted alk(en)yl, alkoxyalkyl, or alkylthioalkyl; Q is a bond, CO, (un)substituted aryl, heteroaryl, or heterocycloalkyl; J is an amino group, including ureido groups; R4 is H, (un)substituted alkyl or arylalkyl; R5 is -M-G-A, where M and A are (un)substituted aryl or heteroaryl; G is a bond, CH2, -alkyl-O-, -O-alkyl-, O, S, SO, or SO2 (with provisos)] useful for inhibiting the ADAM-10 protein, also known as human Kuzbanian. Such compds. are useful in the in vitro study of the role of ADAM-10 (and its inhibition) in biol. processes. Pharmaceutical compns. comprising one or more ADAM-10 inhibitors are useful for the treatment of cancer, arthritis, and diseases related to angiogenesis. The invention also provides methods for making bis-aryl ether sulfonyl chloride intermediates. Thus, claimed compound

N2-[[6-(3-fluorophenyl)pyridin-3-yl]sulfonyl]-N1-hydroxy-D-argininamide showed IC50 < 50 nM for inhibition of ADAM-10.

- OS.CITING REF COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS)
- REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

STRUCTURE SEARCH

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STRUCTURE FILE UPDATES: 13 SEP 2011 HIGHEST RN 1332075-54-3 DICTIONARY FILE UPDATES: 13 SEP 2011 HIGHEST RN 1332075-54-3

CAS Information Use Policies apply and are available at:

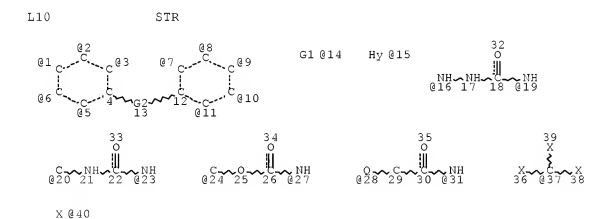
http://www.cas.org/legal/infopolicy.html

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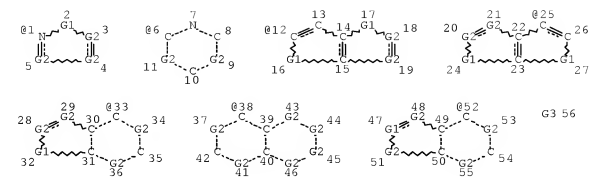
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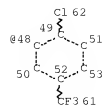
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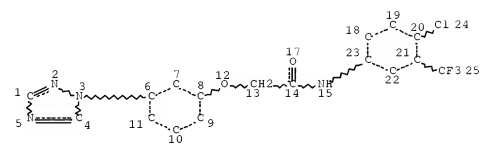
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FILE COVERS 1907 - 14 Sep 2011 VOL 155 ISS 12

FILE LAST UPDATED: 13 Sep 2011 (20110913/ED)

REVISED CLASS FIELDS (/NCL) LAST RELOADED: Jun 2011

USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Jun 2011

CAplus now includes complete International Patent Classification (IPC) reclassification data for the second quarter of 2011.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

'OBI' IS DEFAULT SEARCH FIELD FOR 'CAPLUS' FILE

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L86 77 L84 AND (PD<20030829 OR AD<20030829 OR PRD<20030829)

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L87 84 (L85 OR L86)

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L87 ANSWER 1 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2005:641873 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 143:153299

TITLE: Preparation of substituted urea derivatives for use in

treating heart failure

INVENTOR(S): Morgan, Bradley Paul; Elias, Kathleen A.; Kraynack, Erica Anne; Lu, Pu-Ping; Malik, Fady; Muci, Alex;

Qian, Xiangping; Smith, Whitney Walter; Tochimoto,

Todd; Tomasi, Adam Lewis; Morgans, David J.

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 64 pp., Cont.-in-part of Appl.

No. PCT/US04/001069.

CODEN: USXXCO

DOCUMENT TYPE: %atent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

	PATENT NO.					KIND DATE				APPL	ICAT	ION :	DATE						
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								0805		VO 20	04-U	S106	9		20	0401	14 <		
	WO 2004	10647	30		A3		2005	0324											
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 143:153299

The present invention provides substituted urea derivs., pharmaceutical compns. containing the derivs., and methods for the treatment of heart failure including congestive heart failure, particularly systolic heart failure. The compns. are selective modulators of the cardiac sarcomere, for example, potentiating cardiac myosin. The ureas of the invention are

represented by the formula R1NHC(O)NHR2 wherein: R1 is optionally substituted aryl or heteroaryl; and R2 is optionally substituted aryl, aralkyl; cycloalkyl, heteroaryl, heteroaralkyl or heterocyclyl, including single stereoisomers, mixts. of stereoisomers, and the pharmaceutically acceptable salts, solvates, and solvates of pharmaceutically acceptable salts thereof.

IT 1056969-80-2 1056969-81-3

RL: PRPH (Prophetic)

(Preparation of substituted urea derivatives for use in treating heart failure)

RN 1056969-80-2 CAPLUS

CN Urea, N-[3-fluoro-5-(3-pyridinyloxy)phenyl]-N'-[[4-(1H-imidazol-1-yl)phenyl]methyl]- (CA INDEX NAME)

RN 1056969-81-3 CAPLUS

CN Urea,

 $\begin{tabular}{ll} N-[3-fluoro-5-(3-pyridinyloxy)phenyl]-N'-[[4-(1-methyl-1H-pyrazol-4-yl)phenyl]methyl]- (CA INDEX NAME) \end{tabular}$

PAGE 1-A

PAGE 2-A

OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD

(2 CITINGS)

REFERENCE COUNT: 105 THERE ARE 105 CITED REFERENCES AVAILABLE FOR

THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L87 ANSWER 2 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2005:122807 CAPLUS Full-text

DOCUMENT NUMBER: 142:219152

TITLE: Preparation of tetrahydro-2H-thiopyran-4-carboxamides

as anti-herpesvirus agents

INVENTOR(S): Kontani, Toru; Miyata, Junji; Hamaguchi, Wataru;

Kawano, Tomoaki; Kamikawa, Akio; Suzuki, Hiroshi;

Sudo, Kenji

PATENT ASSIGNEE(S): Yamanouchi Pharmaceutical Co., Ltd., Japan; Rational

Drug Design Laboratories

SOURCE: U.S. Pat. Appl. Publ., 13 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

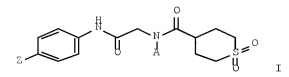
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

US 20050032855	A1	20050210		20040806 <
US 6903125 AU 2004263448	B2	20050607		
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AU 2004263448	B2	20090205		
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KR 2006073928		20060629		20060207 <
MX 2006001526		20060525		20060208 <
US 20060229295	Δ1	20061012		20060208 <
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US 7465748 NO 2006001100	Δ	20061210	NO 2006-1100	20060307 <
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INTOINTIT ALL DIV. TIVE O			WO 2004-JP11573	W 20040805
ASSIGNMENT HISTORY FOR H	S PATEN	IT AWATI.AR		

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): CASREACT 142:219152; MARPAT 142:219152 GI



AB A pharmaceutical drug, particularly a novel compound useful for the prophylaxis or therapeutic treatment of various diseases involving

infections with viruses of the Herpesviridae family, specifically various herpesvirus infections such as varicella (chicken pox) via varicella zoster virus, varicella zoster via recurrent infection with latent varicella zoster virus, herpes labialis and herpes encephalitis via HSV-1 and genital herpes via HSV-2 infection, is disclosed. The title compds. I [\mathbb{Z} =

1,2,4-oxadiazol-3-yl or 4-oxazolyl; A = Ph substituted with one Me group and addnl. with one or two substituents selected from Me, halo; 5-indanyl] have such great anti-virus activity that the oral dosing thereof at a low dose enabled the therapeutic treatment of the diseases. Forty compds. I were prepared Thus, treating Et

{(2,6-dimethylphenyl)[(1,1,-dioxo-tetrahydro-2H-thiopyran-4-

yl)carbonyl]amino}acetate (preparation given) with NaOH solution followed by reacting the crude acid with [4-(1,3-oxazol-4-yl)phenyl]amine afforded I [Z = 1,3-oxazol-4-yl; A = 2,6-(Me)2C6H3] which showed inhibitory activity in HSV-1 infected hairless mice model (93%), and EC50 of 0.075 μ M in anti-VZV activity assay.

IT 841301-36-8P 841301-37-9P 841301-39-1P 841301-40-4P 841301-41-5P 841301-42-6P 841301-43-7P 841301-44-8P 841301-45-9P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of tetrahydro-2H-thiopyran-4-carboxamides as anti-herpesvirus

agents)

RN 841301-36-8 CAPLUS

CN 2H-Thiopyran-4-carboxamide,

N-(4-fluoro-2,6-dimethylphenyl)tetrahydro-N-[2-

[[4-(1,2,4-oxadiazol-3-yl)phenyl]amino]-2-oxoethyl]-, 1,1-dioxide (CA INDEX NAME)

RN 841301-37-9 CAPLUS

CN 2H-Thiopyran-4-carboxamide,

N-(4-fluoro-3-methylphenyl)tetrahydro-N-[2-[[4-

(1,2,4-oxadiazol-3-yl)phenyl]amino]-2-oxoethyl]-, 1,1-dioxide (CA INDEX NAME)

RN 841301-39-1 CAPLUS

CN 2H-Thiopyran-4-carboxamide,

N-(3-bromo-4-methylphenyl)tetrahydro-N-[2-[[4-(1,2,4-oxadiazol-3-yl)phenyl]amino]-2-oxoethyl]-, 1,1-dioxide (CA INDEX NAME)

RN 841301-40-4 CAPLUS

CN 2H-Thiopyran-4-carboxamide,

 $\begin{tabular}{ll} N-(3-fluoro-4-methylphenyl) tetrahydro-N-[2-[[4-(1,2,4-oxadiazol-3-yl)phenyl]amino]-2-oxoethyl]-, 1,1-dioxide (CA INDEX NAME) \\ \end{tabular}$

RN 841301-41-5 CAPLUS

CN 2H-Thiopyran-4-carboxamide,

N-(3-chloro-4-methylphenyl)tetrahydro-N-[2-[[4-(1,2,4-oxadiazol-3-yl)phenyl]amino]-2-oxoethyl]-, 1,1-dioxide (CA INDEX NAME)

RN 841301-42-6 CAPLUS CN 2H-Thiopyran-4-carboxamide,

N-(4-chloro-3-methylphenyl)tetrahydro-N-[2-[[4-(1,2,4-oxadiazol-3-yl)phenyl]amino]-2-oxoethyl]-, 1,1-dioxide (CA INDEX NAME)

RN 841301-43-7 CAPLUS
CN 2H-Thiopyran-4-carboxamide,
N-(4-fluoro-3,5-dimethylphenyl)tetrahydro-N-[2[[4-(1,2,4-oxadiazol-3-yl)phenyl]amino]-2-oxoethyl]-, 1,1-dioxide (CA INDEX NAME)

RN 841301-44-8 CAPLUS
CN 2H-Thiopyran-4-carboxamide,
N-(3-fluoro-2,4-dimethylphenyl)tetrahydro-N-[2[[4-(1,2,4-oxadiazol-3-yl)phenyl]amino]-2-oxoethyl]-, 1,1-dioxide (CA INDEX NAME)

RN 841301-45-9 CAPLUS

CN 2H-Thiopyran-4-carboxamide,

N-(2-fluoro-4-methylphenyl)tetrahydro-N-[2-[[4-

(1,2,4-oxadiazol-3-yl)phenyl]amino]-2-oxoethyl]-, 1,1-dioxide (CA INDEX NAME)

OS.CITING REF COUNT: 6 THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD

(10 CITINGS)

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 3 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2005:120889 CAPLUS Full-text

DOCUMENT NUMBER: 142:198069

TITLE: Preparation of diaryl and arylheteroaryl urea

derivatives, in particular pyrazolylphenyl ureas, as

5-HT2A serotonin receptor modulators

INVENTOR(S): Teegarden, Bradley; Jayakumar, Honnappa; Li, Hongmei;

Strah-Pleynet, Sonja; Dosa, Peter Ian

PATENT ASSIGNEE(S): Arena Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 260 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.					KINI	D :	DATE			APPL:	ICAT	ION I	DA	DATE						
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                                                                               IN 2005-KN2588
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OTHER SOURCE(S): CASREACT 142:198069; MARPAT 142:198069
GΙ
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$$\begin{array}{c} \text{MeO} \\ \text{Me} \\ \text{Me} \\ \text{H} \\ \text{H} \\ \text{H} \\ \text{CF}_{3} \end{array} \qquad \text{II}$$

ΙT

Title compds. I [wherein R1 = (un)substituted hetero/aryl; R2 = H, alk(en/yn)yl, cycloalkyl; R3 = H, alk(en/yn)yl, alkylcarboxamido, alkylsulfonamido, carboxy, cyano, halo, etc.; R4 = NH2, CN, OH, SH, NO2, sulfonamido, halo, alk(en/yn)yl, acyl, acyloxy, etc.; R5 = acyl, acyloxy, alk(en/yn)yl, NH2, halo, haloalkylthio, etc.; R6a, R6b, R6c = independently H, acyl, acyloxy, alk(en/yn)yl, CN, OH, SH, NO2, alkylsulfinyl, alkylureyl, etc.; R7, R8 = independently H, alkyl; X = O, S; Q = a bond, (un)substituted alkylene; and their pharmaceutically acceptable salts, hydrates and solvates] were prepared as modulators of the 5-HT2A serotonin receptor. Thus, reacting

[3-(4-Bromo-2-methyl-2H-pyrazol-3-yl)-4-methoxyphenyl]amine (preparation given) with 4-Chloro-2-(trifluoromethyl)phenyl isocyanate gave II in 60% yield. Nine biol. tests are given. The majority of I showed IC50 activities in the 5-HT2A IP3 accumulation assay of at least about 10 μ M. Selected I attenuated DOI-induced hypolocomotion in rats, demonstrating their inverse agonistic activity. Thus, I and their pharmaceutical compns. thereof are directed to methods useful in the prophylaxis or treatment of platelet aggregation, coronary artery disease, myocardial infarction, transient ischemic attack, angina, stroke, atrial fibrillation, reducing the risk of blood clot formation, asthma or symptoms thereof, agitation or a symptom, behavioral disorders, drug induced psychosis, excitative psychosis, Gilles de la Tourette's syndrome, manic disorder, organic or NOS psychosis, psychotic disorder, psychosis, acute schizophrenia, chronic schizophrenia, NOS schizophrenia and related disorders, and sleep disorders, sleep disorders, diabetic-related disorders and the like. The invention also relates to the method of prophylaxis or treatment of 5-HT2A serotonin receptor mediated disorders in combination with a dopamine D2 receptor antagonist such as haloperidol, administered sep. or together. 839715-19-4P, 1-(4-Chlorobenzyl)-3-[4-(3-dimethylaminopropoxy)-3-

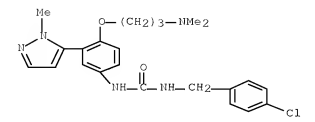
(2-methyl-2H-pyrazol-3-yl)phenyl]urea
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(Uses)

(drug candidate; preparation of pyrazolylphenyl ureas as 5-HT2A serotonin receptor modulators for treatment of heart disease, stroke, psychosis, sleep disorder and other disorders)

RN 839715-19-4 CAPLUS

CN Urea, N-[(4-chlorophenyl)methyl]-N'-[4-[3-(dimethylamino)propoxy]-3-(1-methyl-1H-pyrazol-5-yl)phenyl]- (CA INDEX NAME)



OS.CITING REF COUNT: 7 THERE ARE 7 CAPLUS RECORDS THAT CITE THIS RECORD

(7 CITINGS)

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 4 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2005:34603 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 142:134589

TITLE: Preparation of indazole derivatives for treating or

preventing diseases associated with protein kinases

INVENTOR(S): Bhagwat, Shripad S.; Satoh, Yoshitaka; Sakata, Steven

T.; Buhr, Chris A.; Albers, Ronald; Sapienza, John; Plantevin, Veronique; Chao, Qi; Sahasrabudhe, Kiran;

Ferri, Rachel; Narla, Rama K.

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 240 pp., Cont.-in-part of U.S.

Ser. No. 414,839.

CODEN: USXXCO

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 20050009876	A1	20050113	US 2003-718185	20031119 <
US 20020103229	A1	20020801	US 2001-910950	20010723 <
US 6897231	В2	20050524		
US 20040127536	A1	20040701	US 2003-414839	20030416 <
US 7211594	B2	20070501		
US 20070060616	A1	20070315	US 2006-512836	20060830 <
PRIORITY APPLN. INFO.:			US 2000-221799P	P 20000731 <
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OTHER SOURCE(S): MARPAT 142:134589

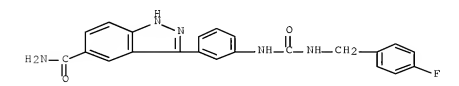
AB Methods of treating or preventing diseases associated with protein kinases, including tyrosine kinases, such as proliferative diseases, inflammatory diseases, abnormal angiogenesis and diseases related thereto, atherosclerosis, macular degeneration, diabetes, obesity, pain and others, comprising administering to a patient in need thereof an effective amount of the title indazole I [A = a direct bond, (CH2)a, (CH2)bCH:CH(CH2)c, or (CH2) bC.tplbond.C(CH2)c; R1 = (un) substituted aryl, heteroaryl or heterocycle fused to Ph; R2 = R3, R4, (CH2)bC(0)R5, (CH2)bC(:0)OR5, (CH2) bC (O) NR5R6, (CH2) bC (O) NR5 (CH2) cC (O) R6, (CH2) bNR5C (O) R6, (CH2) bNR5C(0) NR6R7, (CH2) bNR5R6, (CH2) bOR5, (CH2) bSOdR5 or (CH2) bSO2NR5R6; a = 1-6; b, c = 0-4; d = 0-2; R3 = halo, hydroxy, carboxy, alkyl, alkoxy, haloalkyl, etc.; R4 = (un)substituted alkyl, aryl, arylalkyl, heterocycle or heterocyclealkyl, or R4 = halo or OH; R5-R7 = H, (un) substituted alkyl, aryl, arylalkyl, heterocycle or heterocyclealkyl], are disclosed. Many of the claimed compds. I have IC50 values $\leq 0.5 \mu M$ in the JNK2 assay, e.g. 5-[3-(4-fluorophenyl)-1H-indazol-5-yl]-2H-1,2,3,4-tetrazole. Although the methods of preparation are not claimed, >400 example prepns. are included. ΙT 716322-98-4

RL: PRPH (Prophetic)

(Preparation of indazole derivatives for treating or preventing diseases associated with protein kinases)

RN 716322-98-4 CAPLUS

CN 1H-Indazole-5-carboxamide, 3-[3-[[[[(4-fluorophenyl)methyl]amino]carbonyl]amino]phenyl]- (CA INDEX NAME)



OS.CITING REF COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (5 CITINGS)

L87 ANSWER 5 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2004:902232 CAPLUS Full-text

DOCUMENT NUMBER: 141:374691

TITLE: Anthranilic acid derivatives useful in treating

infection with hepatitis C virus

INVENTOR(S): Bloom, Jonathan D.; Bailey, Thomas R.

PATENT ASSIGNEE(S): Wyeth, John, and Brother Ltd., USA; Viropharma

Incorporated

SOURCE: PCT Int. Appl., 38 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.				KIND DATE			APPLICATION NO.					DATE					
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 141:374691

AB The present invention provides pharmaceutical compns. comprising anthranilic acid derivs. useful in treating hepatitis C infection by virtue of their ability to inhibit hepatitis C polymerase (NS5B). The present invention also provides methods of treating hepatitis C infection by administering to a mammal the pharmaceutical compns. of the present invention.

IT 782481-09-8

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(anthranilic acid derivs. for treatment of hepatitis C virus infection)

RN 782481-09-8 CAPLUS

CN Acetamide, 2-(4-chlorophenoxy)-N-[2-(2H-tetrazol-5-yl)phenyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD

(2 CITINGS)

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 6 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2004:780704 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 141:296035

TITLE: Preparation of oxopyrazolocinnolines as CD80

inhibitors useful as immunomodulators

INVENTOR(S): Mathews, Ian Richard
PATENT ASSIGNEE(S): Avidex Limited, UK
SOURCE: PCT Int. Appl., 76 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 141:296035

Ι

GΙ

Title compds. [I; R1, R3 = H, F, C1, Br, NO2, cyano, alkyl, fluoroalkyl, chloroalkyl, alkoxy, fluoroalkoxy; R4 = CO2H (ester), CONR6R7, NR7COR6, NR7COOR6, NHCONR6R7, NHCSNR6R7; R6 = H, (Alk)mQ; m = 0, 1; Alk = (substituted) alkylene, alkenylene, alkynylene, carbocyclylene which may contain ≥1 O, S, NR8; R8 = H, alkyl, alkenyl, alkynyl, cycloalkyl; Q = H, NR9R10; R9, R10 = H, alkyl, alkenyl, alkynyl, cycloalkyl, ester group, (substituted) carbocyclyl, heterocyclyl; R9R10N = (substituted) heterocyclyl; R7 = H, alkyl; R6R7 = atoms to form (substituted) heterocyclyl; X = bond, (Z)n(Alk), (Alk)(Z)n; Z = O, S, NH; n = 0, 1], were prepared Thus, 4-(3-oxo-1,3-dihydro-2H-pyrazolo[4,3-c]cinnolin-2- yl)benzoic acid (preparation given) was stirred with DMF, diisopropylethylamine, 3-dimethylaminopropylamine, and HTBU at room temperature for 2 h to give 40% N-[(3-dimethylamino)propyl]

 $4\text{-}(3\text{-}oxo\text{-}1,3\text{-}dihydro\text{-}2H\text{-}pyrazolo\text{[}4,3\text{-}c\text{]}cinnolin\text{-}2\text{-}y1\text{)}}$ benzamide (AV1142005). The latter inhibited interleukin-2 production by human Jurkat T cells by 65% at 30 $\mu\text{M}.$

IT 763143-44-8P 763144-94-1P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of oxopyrazolocinnolines as CD80 inhibitors useful as immunomodulators)

RN 763143-44-8 CAPLUS

CN Urea, N-[4-(1,3-dihydro-3-oxo-2H-pyrazolo[4,3-c]cinnolin-2-yl)phenyl]-N'[(4-fluorophenyl)methyl]- (CA INDEX NAME)

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RN 763144-94-1 CAPLUS

CN Urea, N-[4-(1,3-dihydro-3-oxo-2H-pyrazolo[4,3-c]cinnolin-2-yl)phenyl]-N'[(3-fluorophenyl)methyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ \end{array}$$

OS.CITING REF COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD

(7 CITINGS)

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 7 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2004:566609 CAPLUS Full-text.

DOCUMENT NUMBER: 141:123608

TITLE: Preparation of pyrrolopyridinones as mitogen activated

protein kinase-activated protein kinase-2 inhibiting

compounds

INVENTOR(S): Anderson, David R.; Mahoney, Matthew W.; Phillion,

Dennis P.; Rogers, Thomas E.; Meyers, Marvin J.; Poda, Gennadiy; Hegde, Shridhar G.; Singh, Megh; Reitz, David B.; Wu, Kun K.; Buchler, Ingrid P.; Xie, Jin;

Vernier, William F.

PATENT ASSIGNEE(S): Pharmacia Corporation, USA

SOURCE: PCT Int. Appl., 573 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004058762	A1	20040715	WO 2003-US40811	20031219 <

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PRIORITY APPLN. INFO.:
                                            US 2002-434962P
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                                                                 A1 20031219
                                             WO 2003-US40811
                                                                 W 20031219
                         MARPAT 141:123608
OTHER SOURCE(S):
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GΙ

AB The title compds. [I; Z1, Z3, Z4 = C, N; Z2, Z5 = C, N, S, O, and join together with Z1, Z3 and Z4 to form a ring that is selected from a pyrrole, furan, thiophene, oxazole, thiazole, triazole, and imidazole; when either Z2, or Z5 = O or S, it has no substituent group; when Z1-Z5 form an imidazole ring,

Z1 = C and if Z2 and Z5 = N, one is unsubstituted and Z3 and Z4 = C, if Z3and Z5 = N, Z5 is unsubstituted and Z2 and Z4 = C, and if Z2 and Z4 = N, Z2is unsubstituted and Z3 and Z5 = C; when Z1-Z5 form an oxazole or thiazole ring, Z1, Z3 and Z4 = C and one of Z2 and Z5 = N that is unsubstituted; when Z1-Z5 form a triazole ring, Z2 and Z5 = N that is unsubstituted; T = C, N; p = 0-3; X = C, S; Ra = (un) substituted 5-6 membered hetero(aryl) or partially unsatd. 5-6 membered ring; R2, R5, R50-R53, R56 = absent, H, alkyl, aryl, etc.; R54, R55 = oxo, absent] which inhibit mitogen activated protein kinase-activated protein kinase-2 (MK-2), were prepared Thus, reacting 2-(2-chloropyridin-4-y1)-1,5,6,7-tetrahydro-4H-pyrrolo[3,2-c]pyridin-4-o ne (preparation given) with 3-thiopheneboronic acid in the presence of Cs2CO3, Pd(PPh3)4 in DMF afforded 57% II.TFA. The compds. I were tested for MK-2 inhibition activity (biol. data given for over 800 compds). Methods of using compds. I for the inhibition of MK-2, and for the prevention or treatment of a disease or disorder that is mediated by $INF\alpha$, are described, where the method involves administering to the subject an MK-2 inhibiting compound I. Therapeutic compns., pharmaceutical compns. and kits which contain the present MK-2 inhibiting compds. I are also described. [This abstract record is one of 2 records for this document necessitated by the large number of index entries required to fully index the document and publication system constraints.]

IT 724730-57-8P 724730-68-1P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of pyrrolopyridinones as mitogen activated protein kinase-activated protein kinase-2 inhibiting compds. for preventing or treating a $\text{TNF}\alpha$ mediated diseases)

RN 724730-57-8 CAPLUS

CN Acetamide, 2-[2-methoxy-4-[4-(4,5,6,7-tetrahydro-4-oxo-1H-pyrrolo[3,2-c]pyridin-2-y1)-2-pyrimidiny1]phenoxy]-N-[3-(trifluoromethy1)pheny1]-

(CA

INDEX NAME)

RN 724730-68-1 CAPLUS

CN Acetamide, 2-[2-ethoxy-4-[4-(4,5,6,7-tetrahydro-4-oxo-1H-pyrrolo[3,2-c]pyridin-2-yl)-2-pyrimidinyl]phenoxy]-N-[3-(trifluoromethyl)phenyl]-

(CA INDEX NAME)

OS.CITING REF COUNT: 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS)

L87 ANSWER 8 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2004:533982 CAPLUS Full-text

DOCUMENT NUMBER: 141:89085

TITLE: Preparation of indazole derivatives as JNK enzyme

inhibitors

INVENTOR(S): Bhagwat, Shripad S.; Satoh, Yoshitaka; Sakata, Steven

T.; Buhr, Chris A.; Albers, Ronald; Sapienza, John; Plantevin, Veronique; Chao, Qi; Sahasrabudhe, Kiran;

Ferri, Rachel

PATENT ASSIGNEE(S): Signal Pharmaceuticals, LLC, USA

SOURCE: U.S. Pat. Appl. Publ., 275 pp., Cont.-in-part of U.S.

Ser. No. 910,950.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT NO.	KIND D	ATE	APPLICATION NO.	DATE			
US 20040127536		0040701	US 2003-414839	20030416 <			
US 7211594	B2 2	0070501					
US 20020103229	A1 2	0020801	US 2001-910950	20010723 <			
US 6897231	B2 2	0050524					
US 20040077877	A1 20	0040422	US 2003-673121	20030926 <			
US 7220771	B2 2	0070522					
US 20050009876	A1 20	0050113	US 2003-718185	20031119 <			
AU 2004232981	A1 20	0041104	AU 2004-232981	20040416 <			
CA 2522682	A1 2	0041104	CA 2004-2522682	20040416 <			
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OTHER SOURCE(S): MARPAT 141:89085

GΙ

$$\mathbb{R}^2$$

$$\mathbb{A}_{\mathbb{R}^1}$$
 \mathbb{R}^1

AΒ Indazole derivs. I [A = a bond, (CH2)a, (CH2)bCH:CH(CH2)c, (CH2)bC.tplbond.C(CH2)c; R1 = (un)substituted aryl, heteroaryl or heterocycle fused to Ph; R2 = R3, R4, (CH2)bC(0)R5, (CH2)bC(:0)OR5, (CH2) bC (O) NR5R6, (CH2) bC (O) NR5 (CH2) cC (O) R6, (CH2) bNR5C (O) R6, (CH2)bNR5C(0)NR6R7, (CH2)bNR5R6, (CH2)bOR5, (CH2)bSOdR5 or (CH2)bSO2NR5R6; a = 1-6; b, c = 0-4; d = 0-2; R3 = halo, OH, CO2H, carboxy, etc.; R4 = (un) substituted alkyl, aryl, arylalkyl, heterocycle or heterocyclealkyl, or R4 = halo or OH; R5-R7 = H, (un)substituted alkyl, aryl, arylalkyl, heterocycle or heterocyclealkyl; with the provisos] having activity as selective inhibitors of JNK, are disclosed. Such compds. I have utility in the treatment of a wide range of conditions that are responsive to JNK inhibition. Thus, methods of treating such conditions are also disclosed, as are pharmaceutical compns. containing one or more compds. of the above compds. Many of the claimed compds. have IC50 values $\leq 0.5~\mu\mathrm{M}$ in the JNK2 assay, e.g. 5-[3-(4-fluorophenyl)-1H-indazol-5-yl]-2H-1,2,3,4-tetrazole.Although the methods of preparation are not claimed, >400 example prepns. are included.

IT 716322-54-2P 716322-98-4P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of indazole derivs. as JNK enzyme inhibitors)

RN 716322-54-2 CAPLUS

CN Carbamic acid, [3-[5-(aminocarbonyl)-1H-indazol-3-yl]phenyl]-, (4-fluorophenyl)methyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 716322-98-4 CAPLUS

CN 1H-Indazole-5-carboxamide, 3-[3-[[[(4-

fluorophenyl)methyl]amino]carbonyl]amino]phenyl]- (CA INDEX NAME)

$$\begin{array}{c|c} H_{2N-C} & \stackrel{H}{\longrightarrow} N \\ & \stackrel{N}{\longrightarrow} N \\ & \stackrel{O}{\longrightarrow} NH-CH_{2} \\ & \stackrel{F}{\longrightarrow} \end{array}$$

OS.CITING REF COUNT: 32 THERE ARE 32 CAPLUS RECORDS THAT CITE THIS

RECORD (69 CITINGS)

REFERENCE COUNT: 79 THERE ARE 79 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 9 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2004:390211 CAPLUS Full-text

DOCUMENT NUMBER: 140:406638

TITLE: Preparation of arylamides as melanin concentrating

hormone (MCH) receptor antagonists.

INVENTOR(S): Stenkamp, Dirk; Mueller, Stephan Georg; Roth, Gerald

Juergen; Lustenberger, Philipp; Rudolf, Klaus;

Lehmann-Lintz, Thorsten; Arndt, Kirsten; Lotz, Ralf R.

H.; Lenter, Martin; Wieland, Heike-Andrea

PATENT ASSIGNEE(S): Boehringer Ingelheim Pharma GmbH & Co. Kg, Germany; et

al.

SOURCE: PCT Int. Appl., 276 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT NO.			KIND DATE		APPLICATION NO.					DATE							
WO 20040	3976	54		A1	2	20040)513	V	VO 20	03-E	P119	33		20	0310	28 <	-
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OTHER SOURCE(S):
                         MARPAT 140:406638
     R1R2NXYZNR3COWABb [R1, R2 = H, (substituted) alkyl, cycloalkyl,
AΒ
     heterocyclyl, Ph, pyridyl; R1R2 = alkylene optionally interrupted by CH:N,
     CH:CH, O, S, SO, SO2, CO, imino, etc.; R3 = H, alkyl, cycloalkyl,
     cycloalkylalkyl; X = alkylene optionally interrupted by CH:CH, C.tplbond.C,
     O, S, SO, SO2, CO, imino; W = CR6aR6bO, CR7a:CR7c, etc.; Z = bond, (fused)
     (alkyl-substituted) alkylene; Y, A, B = Cy; b = 0, 1; Cy = (substituted)
     (unsatd.) carbocyclyl, Ph, (aromatic) heterocyclyl; R6a, R6b = H, alkyl, CF3;
     R7a, R7c = H, F, Cl, alkyl, CF3; with provisos and specific exceptions], were
     prepared for treatment of obesity, diabetes, heart failure,
     arteriosclerosis, hypertension, arthritis, mastocytosis, depression,
     anxiety, etc. Thus, Me aminoacetate hydrochloride, Et3N, and
     N-[3-chloro-4-(2-oxoethoxy)phenyl]-2-(2,4-dichlorophenoxy)acetamide in
     CH2Cl2/THF were treated with NaBH(OAc)3 followed by stirring for 3 h to give
     78% Me [2-[2-chloro-4-[2-(2,4-
     dichlorophenoxy)acetylamino]phenoxy]ethylamino]acetate. Tested title
     compds. bound to MCH-1 receptors with IC50 = 17-41 nM.
IT
     689299-57-8P
     RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
        (claimed compound; preparation of arylamides as melanin concentrating
hormone (MCH)
        receptor antagonists)
RN
     689299-57-8 CAPLUS
CN
     Acetamide,
N-[3-chloro-4-[2-(dimethylamino)ethoxy]phenyl]-2-[2-chloro-4-(3-
     pyridinyl)phenoxy]- (CA INDEX NAME)
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IT 689302-21-4P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of arylamides as melanin concentrating hormone (MCH) receptor

antagonists)

RN 689302-21-4 CAPLUS

CN Acetamide,

N-[3-chloro-4-[2-(diethylamino)ethoxy]phenyl]-2-[2-chloro-4-(3-pyridinyl)phenoxy]- (CA INDEX NAME)

OS.CITING REF COUNT: 6 THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD

(6 CITINGS)

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 10 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2004:60473 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 140:128423

TITLE: Preparation of heterocyclylbenzoylureas for treating

type 2 diabetes

INVENTOR(S): Schoenafinger, Karl; Defossa, Elisabeth; Kadereit,

Dieter; Von Roedern, Erich; Klabunde, Thomas; Burger,

Hans-Joerg; Herling, Andreas; Wendt, Karl-Ulrich

PATENT ASSIGNEE(S): Aventis Pharma Deutschland GmbH, Germany

SOURCE: PCT Int. Appl., 67 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

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WO 2004007455	A1	20040122	WO 2003-EP7078	20030703 <

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A1 20070125

US 2006-464907

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PRIORITY APPLN. INFO.:
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OTHER SOURCE(S): MARPAT 140:128423
GΙ
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$$\begin{array}{c|c} \mathbb{R}^4 & & \\ & & \mathbb{R}^3 & \mathbb{R}^1 & \mathbb{R}^2 & \mathbb{H}^{\mathsf{et}_{\mathfrak{m}}} \\ & & \mathbb{R}^3 & \mathbb{R}^$$

AB Title compds. [I; R1, R2 = H, (substituted) A, OA, COA, CO2A, AlkCO2H, AlkCO2A; A = alkyl; Alk = alkylene; R3, R4 = F, C1, Br, OH, NO2, CN, (substituted) A, OA, alkenyloxy, alkynyl; R5 = H, F, C1, Br, OH, NO2, CN, (substituted) A, OA, COA, AlkCO2H, AlkCO2A, SO2A, alkenyloxy, alkynyl; X = H, F, C1, Br, OH, NO2, CN, (substituted) A, COA, AlkCO2H, AlkCO2A, SO2A, alkenyl, alkynyl, OA, SO1-2A, NHA, NA2, CO2H, CO2A, CONH2, CONHA, CONA2, SO2NH2, SO2NHA, SO2NA2, NHCOR6; R6 = H, A, cycloalkyl, cycloalkylalkylene, alkenyl, alkynyl, AlkCO2A, AlkCOA, AlkCO2H, AlkCONH2, aryl, Alkaryl, heteroaryl, Alkheteroaryl, heteroarylcarbonyl; het = 4-7 membered (substituted) heterocyclyl, with the exception of pyrrole; m = 1-5; n, p = 0-3], were prepared Thus, 1-(4-amino-3-fluorophenyl)-1H-[1,2,4]triazole (preparation given) and 2-chloro-4,5-difluorobenzoylisocyanate were stirred 30 min in MeCN to give

1-(2-chloro-4,5-difluorobenzoy1)-3-(2-fluor-4-

[1,2,4]triazol-1-ylphenyl)urea. The latter at 10 μ M gave 94% inhibition of activated glycogen phosphorylase.

IT 648916-89-6P

RN

RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(preparation of heterocyclylbenzoylureas for treating type 2 diabetes)

RN 648916-89-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[2-chloro-4-(2H-tetrazol-5-yl)phenyl]amino]carbonyl]-4-fluoro- (CA INDEX NAME)

IT 648916-81-8P 648916-82-9P 648916-83-0P 648916-84-1P 648916-86-3P 648916-87-4P 648916-88-5P 648916-90-9P 648916-91-0P 648916-92-1P 648917-03-7P 648917-04-8P 648917-05-9P 648917-07-1P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of heterocyclylbenzoylureas for treating type 2 diabetes) $648916-81-8\,$ CAPLUS

CN Benzamide, 2-chloro-4,5-difluoro-N-[[[2-fluoro-4-(1H-1,2,4-triazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

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RN 648916-82-9 CAPLUS

CN Benzamide,

2-chloro-4-fluoro-N-[[[2-methoxy-4-(3-methyl-1H-1,2,4-triazol-5-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 648916-83-0 CAPLUS

CN Benzamide, 2-chloro-4,5-difluoro-N-[[[2-methoxy-4-(3-methyl-1H-1,2,4-triazol-5-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 648916-84-1 CAPLUS

CN 1H-1,2,4-Triazole-3-acetic acid, 5-[4-[[[(2-chloro-4-fluorobenzoyl)amino]carbonyl]amino]-3-methoxyphenyl]- (CA INDEX NAME)

RN 648916-86-3 CAPLUS

CN Benzamide, 2-chloro-N-[[[2-chloro-5-(3-methyl-1H-1,2,4-triazol-5-yl)phenyl]amino]carbonyl]-4,5-difluoro- (CA INDEX NAME)

RN 648916-87-4 CAPLUS

CN Benzamide, 2-chloro-N-[[[2-chloro-3-(1H-1,2,4-triazol-5-yl)phenyl]amino]carbonyl]-4,5-difluoro- (CA INDEX NAME)

RN 648916-88-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-(2,5-dihydro-5-oxo-1H-1,2,4-triazol-3-yl)-2-(trifluoromethoxy)phenyl]amino]carbonyl]-4,5-difluoro- (CA INDEX NAME)

RN 648916-90-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[2-chloro-4-(2H-tetrazol-5-yl)phenyl]amino]carbonyl]-4,5-difluoro- (CA INDEX NAME)

RN 648916-91-0 CAPLUS

CN Benzamide, 2-chloro-4-fluoro-N-[[[4-(2H-tetrazol-5-yl)-2-(trifluoromethoxy)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 648916-92-1 CAPLUS

CN Benzamide, 2-chloro-4,5-difluoro-N-[[[4-(2H-tetrazol-5-yl)-2-(trifluoromethoxy)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 648917-03-7 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-(2,5-dihydro-5-oxo-1,2,4-oxadiazol-3-yl)-2-(trifluoromethoxy)phenyl]amino]carbonyl]-4,5-difluoro- (CA INDEX NAME)

RN 648917-04-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-(2,5-dihydro-5-oxo-1,2,4-oxadiazol-3-yl)-2-(trifluoromethoxy)phenyl]amino]carbonyl]-4-fluoro- (CA INDEX NAME)

RN 648917-05-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[2-chloro-4-(2,5-dihydro-5-oxo-1,2,4-oxadiazol-3-yl)phenyl]amino]carbonyl]-4-fluoro- (CA INDEX NAME)

RN 648917-07-1 CAPLUS

CN Benzamide, N-[[[2-(1H-benzimidazol-2-yl)phenyl]amino]carbonyl]-2-chloro-4,5-difluoro- (CA INDEX NAME)

OS.CITING REF COUNT: 9 THERE ARE 9 CAPLUS RECORDS THAT CITE THIS RECORD

(13 CITINGS)

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 11 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2003:971891 CAPLUS Full-text

DOCUMENT NUMBER: 140:13098

TITLE: Pharmaceutically active compounds having a tricyclic

pyrazolotriazolopyrimidine ring structure and methods

of use

INVENTOR(S): Baraldi, Pier Giovanni; Borea, Pier Andrea

PATENT ASSIGNEE(S): King Pharmaceuticals Research & Development, Inc., USA

SOURCE: PCT Int. Appl., 80 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT NO.				KIND DATE				APPLICATION NO.						DATE		
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WO 2003	10145	55		A2	2	20031	1211	V	vo 20	03-U	S173	13		20	0305	30 <
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PRIORITY APPLN. INFO.:
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 140:13098

AB Tricyclic pyrazolotriazolopyrimidines which possess antagonistic activity for adenosine receptors may be useful for modulating biol. function in the nervous, cardiovascular, renal, respiratory and immune systems. General synthetic schemes and examples of formulations for the compds. are presented.

IT 512845-34-0P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(tricyclic pyrazolotriazolopyrimidines with antagonistic activity for adenosine receptors)

RN 512845-34-0 CAPLUS

CN Acetamide, 2-[4-[5-amino-7-(2-phenylethyl)-7H-pyrazolo[4,3-e][1,2,4]triazolo[1,5-c]pyrimidin-2-yl]phenoxy]-N-(4-iodophenyl)- (CA INDEX NAME)

OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD

(4 CITINGS)

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 12 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2003:633320 CAPLUS Full-text DOCUMENT NUMBER: 139:180075

TITLE: Preparation of pyrrolopyrimidines as tyrosine kinase

inhibitors

INVENTOR(S): Hirst, Gavin C.; Calderwood, David; Munschauer,

Rainer; Arnold, Lee D.; Johnston, David N.; Rafferty,

Paul

PATENT ASSIGNEE(S): Abbott GmbH & Co. KG, USA

SOURCE: U.S. Pat. Appl. Publ., 166 pp., Cont.-in-part of Appl.

No. PCT/US99/21560.

CODEN: USXXCO

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 139:180075

GI

AB The title compds. I [A = (un)] substituted 6-membered aromatic ring, 5-6 membered heteroarom. ring; L = O, S, SO, SO2, etc.; G = a direct bond, (CH2)j (wherein j = 1-6), alkenylene, cycloalkylene, oxaalkylene; R1 = alkyl, cycloalkyl, bicycloalkyl, etc.; R2 = H, alkyl, cycloalkyl, halo, etc.; R3 = alkyl, alkenyl, cycloalkyl, etc.] and physiol. acceptable salts and metabolites thereof, are inhibitors of serine/threonine and tyrosine kinase activity. Several of the kinases, whose activity is inhibited by compds. I, are involved in immunol., hyperproliferative, or angiogenic processes. Thus, the compds. I can ameliorate disease states where angiogenesis or endothelial cell hyperproliferation is a factor. These compds. can be used to treat cancer and hyperproliferative disorders, rheumatoid arthritis, disorders of the immune system, transplant rejections and inflammatory disorders. All exemplified compds. I significantly inhibited either FGFR, PDGFR, KDR, Tie-2, Lck, Fyn, Blk, Lyn, or Src at ≤50 µM, and some significantly inhibited cdc2 at ≤50 μM. 546 Example prepns. are included. For example, addition of piperidine to

4-[4-amino-5-(4-phenoxyphenyl)-7H-pyrrolo[2,3-d]pyrimidin-7-yl]cyclohexanone in DCE and AcOH, followed by treatment with Na[(AcO)3BH], workup and chromatog., gave cis- and trans-II. 262442-33-1P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(target compound; preparation of pyrrolopyrimidinamines as protein kinase inhibitors)

RN 262442-33-1 CAPLUS

TΤ

CN Acetamide, N-[4-[4-amino-7-[cis-4-(4-methyl-1-piperazinyl)cyclohexyl]-7H-pyrrolo[2,3-d]pyrimidin-5-yl]-2-methoxyphenyl]-2-(4-chlorophenoxy)- (CA INDEX NAME)

Relative stereochemistry.

PAGE 1-A

OS.CITING REF COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (5 CITINGS)

L87 ANSWER 13 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2003:591153 CAPLUS Full-text

DOCUMENT NUMBER: 139:164789

TITLE: Preparation of phenylpyrazoles as 5-HT2A serotonin

receptor modulators

INVENTOR(S): Teegarden, Bradley; Drouet, Keith; Jayakumar,

Honnappa; Thomsen, William; Maffuid, Paul; Elwell, Katie; Foster, Richard; Lawless, Michael; Liu, Qian;

Smith, Julian; Feichtinger, Konrad

PATENT ASSIGNEE(S): Arena Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 266 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003062206	A2	20030731	WO 2003-US2059	20030123 <

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WO 2003062206
                                20040108
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             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
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             KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
             FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BF,
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     EP 1509505
                         A2
                                20050302
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PRIORITY APPLN. INFO.:
                                            US 2002-386198P
                                                               P 20020123 <--
                                            US 2002-386384P
                                                                P 20020605
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                                            US 2002-401467P
                                                                    20020805
                                                                Ρ
                                            WO 2003-US2059
                                                                    20030123
<--
                         MARPAT 139:164789
OTHER SOURCE(S):
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AB Title compds. I [wherein R1 = H, halo, NR5R6, OH, or OR7; R2 = H, (cyclo)alkyl, or alkenyl; R3 = halo, carboxy, CN, or (un)substituted alkoxycarbonyl, (cyclo)alkyl, alkenyl, alkynyl, or (hetero)aryl; R4 = (cyclo)alkyl or alkenyl; R5 and R6 = independently H or (un)substituted (cyclo)alkyl, alkenyl, aryl(methyl); or NR5R6 = (un)substituted heterocyclyl; R7 = H or alkyl; A = CO, CS, or SO2; B = (NR11)q(CHR12)m(1,2-cyclopropylidene)nQ1 or OQ2; m, n, and q = independently 0-1; R11 and R12 = independently H,

(cyclo) alkyl, or alkenyl; Q = (un) substituted Ph; Q2 = (un) substituted (cyclo)alkyl, alkenyl, alkynyl, alkylaryl, or aryl(alkyl); and pharmaceutically acceptable salts thereof] were prepared as modulators of the 5-HT2A serotonin receptor. For example, reaction of triphosgene with 3-(3-aminophenyl)-4-bromo-2-methylpyrazole in the presence of TEA in CH2Cl2, followed by addition of 4-(trifluoromethoxy)benzylamine provided the N-(pyrazolylphenyl)urea II (68%). The latter exhibited IC50 values of 1.2 μ M, 0.45 μ M, and 0.0171 μ M for AP-1, WT 5-HT2A, and AP-3, resp., in a competitive binding assay. A number of the compds. of the invention evidenced inverse agonist activity against AP-1 (data given). Thus, I and pharmaceutical compns. thereof are directed to methods useful in the prophylaxis or treatment of reducing platelet aggregation, coronary artery disease, myocardial infarction, transient ischemic attack, angina, stroke, atrial fibrillation, reducing the risk of blood clot formation, asthma or symptoms thereof, agitation or a symptom, behavioral disorders, drug induced psychosis, excitative psychosis, Gilles de la Tourette's syndrome, manic disorder, organic or NOS psychosis, psychotic disorder, psychosis, acute schizophrenia, chronic schizophrenia and NOS schizophrenia, and related disorders (no data). The present invention also relates to the method of prophylaxis or treatment of 5-HT2A serotonin receptor mediated disorders in combination with a dopamine D2 receptor antagonist such as haloperidol, administered sep. or together.

IT 573711-73-6P, 1-[3-(4-Bromo-2-methyl-2H-pyrazol-3-yl)phenyl]-3[(4-fluorophenyl)methyl]urea

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(5-HT2A receptor modulator; preparation of phenylpyrazoles as 5-HT2A serotonin receptor modulators for treatment of heart disease, stroke, psychosis, and other disorders)

RN 573711-73-6 CAPLUS

CN Urea, N-[3-(4-bromo-1-methyl-1H-pyrazol-5-yl)phenyl]-N'-[(4-fluorophenyl)methyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 7 THERE ARE 7 CAPLUS RECORDS THAT CITE THIS RECORD (8 CITINGS)

L87 ANSWER 14 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2003:334911 CAPLUS Full-text

DOCUMENT NUMBER: 138:354000

TITLE: Preparation of dihydroxypyrimidine carboxamide

inhibitors of HIV integrase

INVENTOR(S): Di Francesco, Maria Emilia; Gardelli, Cristina;

Harper, Steven; Matassa, Victor Giulio; Muraglia, Ester; Nizi, Emanuela; Pace, Paola; Pacini, Barbara;

Petrocchi, Alessia; Poma, Marco; Summa, Vincenzo

PATENT ASSIGNEE(S): Istituto Di Ricerche Di Biologia Molecolare P.

Angeletti Spa, Italy

SOURCE: PCT Int. Appl., 315 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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US	7232	819			В2		2007	0619									
US	20070	0830	45		A1	2	20070	1412	Ţ	JS 20	06-5	1683	1		20	0609	07 <
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									1	WO 2	002-	GB47	42	Ī	₩ 2	0021	021
-																	
									1	US 2	004-	4932	79	1	A3 2	0040	420
SIGNM	GNMENT HISTORY FOR US PATENT AVAILABLE									N LS	US D	ISPL	AY F	ORMA'	Τ		

R1 N R4

GI

OTHER SOURCE(S): MARPAT 138:354000

AΒ The title 4,5-dihydroxypyrimidine-6-carboxamides [I; R1 = H, alkyl, haloalkyl, alkoxy, etc.; R2 = H, alkyl, haloalkyl, hydroxyalkyl, etc.; R3 = H, alkyl; R4 = H, alkyl, haloalkyl, etc.] which are inhibitors of HIV integrase and inhibitors of HIV replication, and therefore are useful in the prevention and treatment of infection by HIV and in the prevention, delay in the onset, and treatment of AIDS, were prepared Thus, refluxing N-hydroxythiophene-2-carboximidamide with di-Me acetylenedicarboxylate in CHCl3 followed by reacting the resulting Me 5,6-dihydroxy-2-(2-thienyl)pyrimidine-4-carboxylate with 4-fluorobenzylamine in DMF afforded I [R1 = 2-thienyl; R2 = H; R3 = 4-FC6H4CH2; R4 = H]. The compds. I are employed against HIV infection and AIDS as compds. per se or in the form of pharmaceutically acceptable salts. The compds. I and their salts can be employed as ingredients in pharmaceutical compns., optionally in combination with other antivirals, immunomodulators, antibiotics or vaccines.

IT 519023-70-2P 519023-73-5P 519024-20-5P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(preparation of dihydroxypyrimidine carboxamide inhibitors of HIV integrase)

RN 519023-70-2 CAPLUS

CN 4-Pyrimidinecarboxamide, 2-[2-[[[(2,3-

dichlorophenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6oxo-N-(phenylmethyl)- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ \text{Ph-CH2-NH-C} & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ \end{array}$$

RN 519023-73-5 CAPLUS

CN 4-Pyrimidinecarboxamide, 2-[2-[[[(2,5-

dichlorophenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6oxo-N-(phenylmethyl)- (CA INDEX NAME)

$$\begin{array}{c} \text{C1} \\ \text{NH-CH}_2 - \text{NH-C} \\ \text{HO} \\ \end{array}$$

RN 519024-20-5 CAPLUS

CN 4-Pyrimidinecarboxamide, 2-[3-[[[[(3,4-

dichlorophenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6oxo-N-(phenylmethyl)- (CA INDEX NAME)

OS.CITING REF COUNT: 22 THERE ARE 22 CAPLUS RECORDS THAT CITE THIS

RECORD (31 CITINGS)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 15 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2003:254172 CAPLUS Full-text

DOCUMENT NUMBER: 138:281081

TITLE: Drug screening with non-endogenous, constitutively

activated human serotonin receptors and small molecule

modulators thereof

INVENTOR(S): Behan, Dominic P.; Chalmers, Derek T.; Liaw, Chen W.;

Russo, Joseph F.; Thomsen, William J.

PATENT ASSIGNEE(S): Arena Pharmaceuticals, Inc., USA

SOURCE: U.S., 62 pp., Cont.-in-part of U.S. Ser. No. 60,188.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 17

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6541209	В1	20030401	US 1999-292072	19990414 <
US 6140509	A	20001031	US 1999-292069	19990414 <
US 6420541	B1	20020716	US 2000-767013	20001222 <

US 20030224442	A1	20031204	US 2002-55555	20020123 <
US 20030153004	A1	20030814	US 2002-176255	20020619 <
US 6846919	В2	20050125		
AU 2004203102	A1	20040729	AU 2004-203102	20040708
AU 2004203102	В2	20071018		
US 20050119182	A1	20050602	US 2004-980560	20041102 <
US 7368539	В2	20080506		
AU 2007202139	A1	20070531	AU 2007-202139	20070510
AU 2007202139	B2	20090521		
AU 2007202155	A1	20070607	AU 2007-202155	20070510
AU 2007202155	B2	20090507		
AU 2007202121	A1	20070607	AU 2007-202121	20070511
AU 2007202241	A1	20070607	AU 2007-202241	20070511
AU 2007216751	A1	20071004	AU 2007-216751	20070912 <
AU 2007216752	A1	20071004	AU 2007-216752	20070912 <
AU 2008200231	A1	20080207	AU 2008-200231	20080116
US 20090076254	A1	20090319	US 2008-51754	20080319 <
US 7754866	В2	20100713		
PRIORITY APPLN. INFO.:			US 1997-839449	B2 19970414 <
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			US 1998-112909P	P 19981218
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			US 1999-123000P	P 19990305
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			US 1999-292072	A3 19990414 <
			US 2000-767013	A3 20001222 <
			AU 2002-219890	A3 20011126 <
			US 2002-176255	A1 20020619 <
			AU 2004-202147	A3 20040512
			AU 2004-202476	A3 20040603
			AU 2004-203102	A3 20040708
ACCIONNENT MICEORY BOD M	a 5.5		US 2004-980560	A3 20041102

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

- AB Disclosed herein are non-endogenous, constitutively activated forms of the human 5-HT2A and human 5-HT2C receptors and uses of such receptors to screen candidate compds. Further disclosed herein are candidate compds. identified by the screening method which act at the 5HT2A receptors. Yet further disclosed is a new class of compds. which act at the 5HT2A receptors. IT 247038-30-8P
 - RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(drug screening with non-endogenous, constitutively activated human serotonin receptors and small mol. modulators thereof)

- RN 247038-30-8 CAPLUS
- CN Urea, N-[3-(4-bromo-1-methyl-1H-pyrazol-3-yl)phenyl]-N'-[(4-fluorophenyl)methyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 10 THERE ARE 10 CAPLUS RECORDS THAT CITE THIS

RECORD (11 CITINGS)

REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 16 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2003:221693 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 138:238197

TITLE: Preparation of furo- and thienopyrimidines as TIE-2

and/or VEGFR-2 kinase inhibitors useful against

hyperproliferative diseases

INVENTOR(S): Adams, Jerry Leroy; Bryan, Deborah Lynne; Feng,

Yanhong; Matsunaga, Shinichiro; Maeda, Yutaka;

Miyazaki, Yasushi; Nakano, Masato; Rocher,

Jean-Philippe; Sato, Hideyuki; Semones, Marcus; Silva,

Domingos J.; Tang, Jun

PATENT ASSIGNEE(S): Glaxosmithkline K.K., Japan; Smithkline Beecham

Corporation

SOURCE: PCT Int. Appl., 265 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT	NO.			KIN	D	DATE	ATE APPLICATION NO. DATE										
WO 20030						20030320 WO 2002-US28650 20031127							20020910 <				
W:	ΑE,	AG,	AL,	AM,	AT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
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	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	
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AU 20023	33352	24		A1	2	20030	324	AU 2002-333524						20	0209	10 <	
EP 14252	284			A2	2	20040	0609	F	EP 20	02-7	9818	1		20020910 <			
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JP 2005	50890) 4		Τ	2	20050	0407	Ċ	JP 20	03 - 5	2692	6		20020910 <			
US 2005	00041	142		A1	2	20050	106	US 2004-489052 20040309					09 <				
US 7427	623			В2		20080923											
US 20080	02874	166		A1	2	20081	1120	20 US 2008-169800 20080709 <							09 <		

PRIORITY APPLN. INFO.:

US 2001-318766P P 20010911 <--WO 2002-US28650 W 20020910

<--

US 2004-489052 A3 20040309

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 138:238197

GΙ

AB Furo- and thienopyrimidine derivs. (shown as I; variables defined below; e.g. 4-Amino-3-(4-methoxyphenyl)-2-[3-(methylsulfonylamino)phenyl]furo[2,3-d]pyrimidine), which are useful as TIE-2 (tyrosine kinase containing immunoglobin and EGF homol. domains) and/or VEGFR-2 kinase inhibitors against hyperproliferative diseases are described herein. Enzyme inhibitions by .apprx.60 examples of I are included as ranges; also,

4-amino-3-[4-[[2-fluoro-5-(trifluoromethyl)phenyl]aminocarbonylamino]phenyl]thieno[2,3-d]pyrimidin e exhibited IC50 = $0.0018 \mu M$ in the TIE-2 fluorescence polarization kinase activity assay. For I: X is O or S; A is H, halo, C1-C6 alkyl, aryl, heteroaryl, aryl or heteroaryl substituted with ≥1 R3, heterocyclyl, -RR3, -C(0)OR4, -C(0)NR5R6, -C(0)R4; D is H, halo, C1-C6 alkyl, aryl, heteroaryl, arvl or heteroarvl substituted with ≥1 R3, heterocyclvl, -RR3, -C(0)OR4, -C(O)NR5R6, or -C(O)R4. R is C1-C6 alkylene, C3-C7 cycloalkylene, C1-C6 alkenylene, or C1-C6 alkynylene; R1 is H, C1-C6 alkyl, C1-C6 alkoxy, -SR4, -S(0)2R4, -NR7R7, -NR'N R'''R'''', -N(H)RR3, -C(O)OR7, or -C(O)NR7R7. R2 is H, -OH, -NR7R7 or :NH; R3 is halo, C1-C6 alkyl, C1-C6 haloalkyl, C1-C6 alkoxy, C3-C7 cycloalkoxy, C1-C6 haloalkoxy, aryl, aralkyl, aryloxy, heteroaryl, heterocyclyl, -CN, -NHC(O)R4, -N(R8)HC(O)R4, -NHC(S)R4, -NR5R6, -RNR5R6, -SR4, -S(0)2R4, -RC(0)0R4, -C(0)0R4, -C(0)R4, -C(0)NR5R6, -NHS(O)2R4, -N(S(O)2R4)S(O)2R4, -S(O)2NR5R6, or -NHC(:NH)R4. R4 is H, C1-C6 alkyl, aryl, heteroaryl, heterocyclyl, -RR3, -NR'''R'''', or -NR'NR'''R''''; R5 is H, C1-C6 alkyl, C3-C7 cycloalkyl, cyanoalkyl, -R'R'', aryl, aralkyl, heteroaryl, -NHC(0)OR''', -R'NHC(0)OR''', -R'NHC(O)NR'''R'''', or -R'C(O)OR'''. R6 is H, C1-C6 alkyl, C3-C7 cycloalkyl, cyanoalkyl, -R'R'', aryl, aralkyl, heteroaryl, -C(0)OR''', or -R'C(O)NR'''R'''; R7 is H, C1-C6 alkyl, aryl, or -C(O)OR'''; R8 is C1-C3 alkyl; R' is C1-C3 alkylene; R'' is heteroalkyl or NRR'''R''''; R''' is H, C1-C6 alkyl, aryl, aralkyl, heteroaryl, or C3-C7 cycloalkyl; R'''' is H, C1-C6 alkyl, aryl, heteroaryl, or C3-C7 cycloalkyl. Although the methods of preparation are not claimed, several example prepns. of I are included and characterization data is given for .apprx.480 examples of I.

(drug candidate; preparation of furo- and thienopyrimidines as ${\tt TIE-2}$ and/or

VEGFR-2 kinase inhibitors useful against hyperproliferative diseases)

RN 501695-83-6 CAPLUS

CN Urea, N-[[4-[4-amino-6-(4-methoxyphenyl)furo[2,3-d]pyrimidin-5-yl]phenyl]-N'-[2-fluoro-5-(trifluoromethyl)phenyl]- (CA INDEX NAME)

$$\begin{array}{c} \text{OMe} \\ \text{OMe} \\ \text{OH}_2 \\ \text{OH}_2 \\ \text{OH}_2 \\ \text{OH}_2 \\ \text{OH}_3 \\ \text{OH}_4 \\ \text{OH}_4 \\ \text{OH}_5 \\ \text{OH}_5 \\ \text{OH}_6 \\ \text$$

OS.CITING REF COUNT: 14 THERE ARE 14 CAPLUS RECORDS THAT CITE THIS

RECORD (15 CITINGS)

REFERENCE COUNT: 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 17 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2003:153400 CAPLUS Full-text

DOCUMENT NUMBER: 138:187776

TITLE: Preparation of oxadiazolyl and thiadiazolyl benzoyl

ureas as pesticides

INVENTOR(S): Maurer, Fritz; Erdelen, Christoph; Reckmann, Udo

PATENT ASSIGNEE(S): Bayer CropScience AG, Germany

SOURCE: Ger. Offen., 20 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT	NO.			KIN	D :	DATE			APPL	ICATION NO. DATE						
DE 10139		33		A1 A1	_								20010813 <			
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	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TN,	TR,	TT,	TZ,
	UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZM,	ZW							
RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AT,	BE,	BG,
	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,
	PT,	SE,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	G₩,	ML,	MR,
	NE,	SN,	TD,	ΤG												
AU 20023	32592	29		A1	2	20030	0303	Z	AU 20	02 - 3	2592	9		20020801 <-		
IN 20021	4U006	574		A	2	20050	318	B IN 2002-MU674 200208					05 <			

PRIORITY APPLN. INFO.:

DE 2001-10139721 WO 2002-EP8572

A 20010813 <--W 20020801

<--

OTHER SOURCE(S): MARPAT 138:187776

GI

Title compds. [I; R1 = H, halo; R2 = halo; R3 = halo, (halo)alkyl; n = 0-2; Q = Q1, Q2, Q3; R4 = H, alkyl, alkoxyalkyl, alkoxycarbonylalkyl, alkylcarbonyloxyalkyl, (substituted) aryl, arylalkyl; X = 0, S], were prepared Thus, 4-(5-ethyl-1,3,4-oxadiazol-2-yl)aniline (preparation given) was dropwise treated with 2,6-difluorobenzoyl isocyanate in MeCN at room temperature followed by stirring for 18 h at room temperature to give 86%

N-(2,6-difluorobenzoy1)-N'-[4-(5-ethyl-1,3,4-oxadiazol-2-y1)phenyl]urea. Several I at 0.05% gave 100% kill of Heliothis virescens caterpillars on Glycine max after 7 days.

IT 1053718-64-1 1053718-65-2 1053718-66-3 1053718-69-6 1053718-71-0 1053718-80-1 1053718-82-3 1053718-84-5 1053718-85-6

1053718-99-2

RL: PRPH (Prophetic)

(Preparation of oxadiazolyl and thiadiazolyl benzoyl ureas as pesticides)

RN 1053718-64-1 CAPLUS

CN Benzamide, N-[[[2,5-dichloro-4-[5-(1,1-dimethylethyl)-1,2,4-oxadiazol-3-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

CN Benzamide, 2-chloro-N-[[[2,5-dichloro-4-[5-(1,1-dimethylethyl)-1,2,4-oxadiazol-3-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 1053718-66-3 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[5-(1,1-dimethylethyl)-1,2,4-oxadiazol-3-yl]-2,3-difluorophenyl]amino]carbonyl]- (CA INDEX NAME)

RN 1053718-69-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[2,3-dichloro-4-[5-(1,1-dimethylethyl)-1,2,4-oxadiazol-3-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 1053718-71-0 CAPLUS

CN Benzamide, N-[[[2,3-dichloro-4-[5-(1,1-dimethylethyl)-1,2,4-oxadiazol-3-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 1053718-80-1 CAPLUS

CN Benzamide, N-[[[2,5-dichloro-4-[3-(1,1-dimethylethyl)-1,2,4-oxadiazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$t-Bu = NH - U - NH$$

RN 1053718-82-3 CAPLUS

CN Benzamide, 2-chloro-N-[[[2,5-dichloro-4-[3-(1,1-dimethylethyl)-1,2,4-oxadiazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 1053718-84-5 CAPLUS

CN Benzamide, N-[[[4-[3-(1,1-dimethylethyl)-1,2,4-oxadiazol-5-yl]-2,5-difluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 1053718-85-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[3-(1,1-dimethylethyl)-1,2,4-oxadiazol-5-yl]-2,5-difluorophenyl]amino]carbonyl]- (CA INDEX NAME)

RN 1053718-99-2 CAPLUS

CN Benzamide, N-[[[4-[5-(1,1-dimethylethyl)-1,2,4-oxadiazol-3-yl]-2,3-difluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

IT 498547-80-1P 498547-82-3P 498547-84-5P 498547-85-6P 498547-86-7P 498547-87-8P

498547-88-9P 498547-95-8P

RL: AGR (Agricultural use); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of oxadiazolyl and thiadiazolyl benzoyl ureas as pesticides) 498547 - 80 - 1 $\,$ CAPLUS

CN Benzamide,

RN

N-[[[4-(5-ethyl-1,2,4-oxadiazol-3-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 498547-82-3 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[3-(1,1-dimethylethyl)-1,2,4-oxadiazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 498547-84-5 CAPLUS

CN Benzamide, 2,6-difluoro-N-[[[4-(5-phenyl-1,2,4-oxadiazol-3-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 498547-85-6 CAPLUS

CN Benzamide, N-[[[4-[5-(1,1-dimethylethyl)-1,2,4-oxadiazol-3-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 498547-86-7 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[5-(1,1-dimethylethyl)-1,2,4-oxadiazol-3-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 498547-87-8 CAPLUS

CN Benzamide, N-[[[4-[5-(1,1-dimethylethyl)-1,2,4-oxadiazol-3-yl]-2,5-difluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 498547-88-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[5-(1,1-dimethylethyl)-1,2,4-oxadiazol-3-yl]-2,5-difluorophenyl]amino]carbonyl]- (CA INDEX NAME)

RN 498547-95-8 CAPLUS

CN Benzamide, N-[[[4-[3-(1,1-dimethylethyl)-1,2,4-oxadiazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

L87 ANSWER 18 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2003:151162 CAPLUS Full-text

DOCUMENT NUMBER: 138:321211

TITLE: Design, Synthesis, and Biological Evaluation of C9-

and C2-Substituted

Pyrazolo[4,3-e]-1,2,4-triazolo[1,5-c]pyrimidines as

New A2A and A3 Adenosine Receptor Antagonists

AUTHOR(S): Baraldi, Pier Giovanni; Fruttarolo, Francesca;

Tabrizi, Mojgan Aghazadeh; Preti, Delia; Romagnoli, Romeo; El-Kashef, Hussein; Moorman, Allan; Varani, Katia; Gessi, Stefania; Merighi, Stefania; Borea, Pier

Andrea

CORPORATE SOURCE: Dipartimento di Scienze Farmaceutiche and Dipartimento

di Medicina Clinica e Sperimentale-Sezione di

Farmacologia, Universita di Ferrara, Ferrara, 44100,

Italy

SOURCE: Journal of Medicinal Chemistry (2003), 46(7),

1229-1241

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 138:321211

GΙ

AB Pyrazolo[4,3-e]-1,2,4-triazolo[1,5-c]pyrimidines such as I are prepared as selective adenosine A2a and A3 receptor antagonists.

Pyrazolo[4,3-e]-1,2,4-triazolo[1,5-c]pyrimidines substituted at the 9-position retain receptor affinity but lose selectivity for the adenosine A2a and A3 receptors over other adenosine receptors. Replacement of the furan moiety present in the pyrazolo[4,3-e]-1,2,4-triazolo[1,5-

c]pyrimidine with a Ph or a substituted aromatic ring abolishes affinity at all the adenosine receptor subtypes, demonstrating that the furanyl ring is a necessary structural element to guarantee interaction with the adenosine receptor surface; replacement of the furan ring with an

ortho-ethoxy-substituted aromatic ring did not enhance affinity.

Introduction of a N-methylpiperazinomethyl or morpholinomethyl function at the 5' position of the furanyl ring of I or introduction of a methylsulfanyl moiety at the 9-position of

pyrazolo[4,3-e]-1,2,4-triazolo[1,5-c]pyrimidines yields inhibitors with improved water solubilities but reduced affinities for adenosine A2a and A3 receptors.

IT 512845-34-0P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and structure-activity relationships of

pyrazolo[4,3-e]-1,2,4-triazolo[1,5-c]pyrimidines as potential selective adenosine A2a and A3 receptor antagonists)

RN 512845-34-0 CAPLUS

CN Acetamide, 2-[4-[5-amino-7-(2-phenylethyl)-7H-pyrazolo[4,3-e][1,2,4]triazolo[1,5-c]pyrimidin-2-yl]phenoxy]-N-(4-iodophenyl)- (CA INDEX NAME)

OS.CITING REF COUNT: 50 THERE ARE 50 CAPLUS RECORDS THAT CITE THIS RECORD (51 CITINGS)

REFERENCE COUNT: 29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 19 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2002:925396 CAPLUS Fuli-text

DOCUMENT NUMBER: 138:17998

TITLE: Photothermographic recording paper comprising

polymerizable compound

INVENTOR(S): Takashima, Masanobu; Sato, Hiroshi; Arai, Yoshimitsu;

Hanasaki, Kyoko

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan

SOURCE: Eur. Pat. Appl., 80 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATE		KIND DATE			APPLICATION NO.						DATE						
														_			
EP 12	EP 1262828			A1	2	20021204 EP 2002-11662							20020531 <				
:	R: AT	, BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PΤ	,
	ΙE	, SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR							
JP 20	03050	142		Α	2	2003	0221	Ċ	JP 20	02-9	8759			20	0204	101	<
US 20	0030073	3025		A1	2	2003	0417	Ţ	JS 20	02-1	5821	0		20	0205	31	<
US 6	720124			В2		2004	0413										
PRIORITY APPLN. INFO.:								J	JP 20	01-1	6673	1	A	20	0106	01	<
									JP 2	002-	98759	9	I	4 2	0020	401	

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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 138:17998

AB A recording photothermog. paper includes a support having disposed thereon a recording layer comprising at least: a diazo compound having no diazonio group, as further described in the claims; a coupler compound that colors by reacting with the diazo compound; and a polymerizable compound Since the photothermog. paper according to the present invention contains a polymerizable compound in the recording layer, it shows excellent image fastness, high sensitivity and fixing speed. A highly sensitive image can be recorded and photofixed at high speed using not only UV light but visible to IR light in a completely dry processing system that does not require a developing solution and therefore does not generate waste; there is excellent decoloring at non-image portions (background portions); and a sharp high-contrast black-and-white or color image can be formed.

IT 477704-73-7

RL: TEM (Technical or engineered material use); USES (Uses) (coupler; photothermog. recording paper comprising polymerizable thermoplasticizer compound and coupler that colors by reacting with diazo compound)

RN 477704-73-7 CAPLUS

CN Benzoic acid, 3-chloro-4-[2-[[4-[6-(1,1-dimethylethyl)-3H-pyrazolo[1,5-b][1,2,4]triazol-2-yl]phenyl]amino]-2-oxoethoxy]-, 6-[(2-methyl-1-oxo-2-propen-1-yl)oxy]hexyl ester (CA INDEX NAME)

PAGE 1-B

OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD

(4 CITINGS)

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 20 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2002:770129 CAPLUS Full-text

DOCUMENT NUMBER: 137:279184

TITLE: Preparation of 3-(hetero)aryl pyrazoles with

4,5(3,4)-bicyclic ring fusion as protein kinase

inhibitors

INVENTOR(S): Doyle, Kevin J.; Rafferty, Paul; Steele, Robert W.;

Wilkins, David J.; Arnold, Lee D.; Hockley, Michael;

Ericsson, Anna M.; Iwasaki, Nobuhiko; Ogawa, Nobuo

PATENT ASSIGNEE(S): BASF Aktiengesellschaft, Germany

SOURCE: U.S., 69 pp., Cont.-in-part of WO 2000 27,822.

CODEN: USXXAM

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6462036 WO 2000027822	B1 A2	20021008 20000518	US 2000-573366 WO 1999-US26105	20000517 < 19991104 <
CZ, DE, IN, IS,	DK, DM, EE JP, KE, KG	E, ES, FI, G, KP, KR,	BB, BG, BR, BY, CA, GB, GD, GE, GH, GM, KZ, LC, LK, LR, LS, NZ, PL, PT, RO, RU,	HR, HU, ID, IL, LT, LU, LV, MA,
SK, SL, RW: GH, GM, DK, ES,	TJ, TM, TR KE, LS, MW FI, FR, GB	R, TT, TZ, I, SD, SL, B, GR, IE,	UA, UG, US, UZ, VN, SZ, TZ, UG, ZW, AT, IT, LU, MC, NL, PT, MR, NE, SN, TD, TG	YU, ZA, ZW BE, CH, CY, DE,
CA 2409225 WO 2001087846 WO 2001087846	A1 A2 A3		CA 2001-2409225 WO 2001-US16153	20010517 < 20010517 <

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             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT,
             RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US,
             UZ, VN, YU, ZA, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
     EP 1289525
                          A2
                                20030312
                                           EP 2001-937553
                                                                   20010517 <--
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
     JP 2003533514
                          Τ
                                20031111
                                            JP 2001-584242
                                                                   20010517 <--
    MX 2002011320
                          Α
                                20040910
                                            MX 2002-11320
                                                                   20021115 <--
PRIORITY APPLN. INFO.:
                                            US 1998-107467P
                                                                P 19981106 <--
                                            WO 1999-US26105
                                                                A2 19991104 <--
                                            US 2000-573366
                                                                 A 20000517
<--
                                            WO 2001-US16153
                                                                 W
                                                                    20010517
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
                        CASREACT 137:279184; MARPAT 137:279184
```

Ι

OTHER SOURCE(S):

GI

AB Title compds. I [m = 1-10; X = alkyl, CO, O, oximino, etc.; B = alkyl,cycloalkyl, aryl, pyridyl, thienyl, furyl, pyrrolyl; R1 = H, halo, hydroxy, nitro, cyano, hydroxyamidino, etc.; A = (un) substituted with one or more substituents selected from halo, alkyl, etc.] were prepared For instance, indan-1-one hydrazone (preparation given) was reacted with Me 3, 4, 5-trimethoxybenzoate (THF, n-BuLi, 0°) and subsequently acidified with HC1 (3 M) and heated to reflux for 1 h to give II. I are inhibitors of protein

kinase activity and used for the treatment of, e.g., cancer, diabetic retinopathy, etc.

IT 268563-63-9P, N-(2,5-Difluorobenzyl)-N'-[4-(1,4-

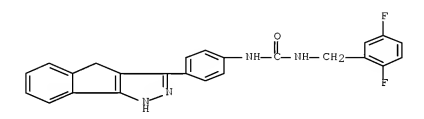
dihydroindeno[1,2-c]pyrazol-3-yl)phenyl]urea

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(kinase inhibitor; 3-(hetero) aryl pyrazoles with 4,5(3,4)-bicyclic ring fusion as protein kinase inhibitors)

RN 268563-63-9 CAPLUS

CN Urea, N-[(2,5-difluorophenyl)methyl]-N'-[4-(1,4-dihydroindeno[1,2-c]pyrazol-3-yl)phenyl]- (CA INDEX NAME)



OS.CITING REF COUNT: 9 THERE ARE 9 CAPLUS RECORDS THAT CITE THIS RECORD

(9 CITINGS)

REFERENCE COUNT: 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 21 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2002:754342 CAPLUS Full-text

DOCUMENT NUMBER: 137:263068

TITLE: Preparation of aryl and biaryl derivatives having

Melanin-concentrating hormone modulatory activity

INVENTOR(S): Hobbs, Douglas W.; Guo, Tao; Hunter, Rachael C.; Gu,

Huizhong

PATENT ASSIGNEE(S): Pharmacopeia, Inc., USA

SOURCE: PCT Int. Appl., 180 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE	E APPL	ICATION NO.	DATE
WO 2002076929	A1 2002	21003 WO 20	02-US8300	20020319 <
W: AE, AG, AL	, AM, AT, AU,	, AZ, BA, BB,	BG, BR, BY,	BZ, CA, CH, CN,
CO, CR, CZ	, DE, DK, DM,	, DZ, EC, EE,	ES, FI, GB,	GD, GE, HR, HU,
ID, IL, IN	, IS, JP, KG,	, KR, KZ, LC,	LK, LR, LT,	LU, LV, MA, MD,
MG, MK, MN	, MX, MZ, NO,	, NZ, PH, PL,	PT, RO, RU,	SE, SG, SI, SK,
SL, TJ, TM	, TN, TR, TT,	, TZ, UA, UZ,	VN, YU, ZA,	ZM
RW: GH, GM, KE	LS, MW, MZ,	, SD, SL, SZ,	TZ, UG, ZM,	ZW, AT, BE, CH,
CY, DE, DK	, ES, FI, FR,	, GB, GR, IE,	IT, LU, MC,	NL, PT, SE, TR,
BF, BJ, CF,	CG, CI, CM,	, GA, GN, GQ,	GW, ML, MR,	NE, SN, TD, TG

CA 2441235	A1	20021003	CA 2002-2441235	20020319 <
CA 2441235	С	20110524		
AU 2002247367	A1	20021008	AU 2002-247367	20020319 <
AU 2002247367	B2	20051027		
US 20030092715	A1	20030515	US 2002-101136	20020319 <
US 7034056	B2	20060425		
EP 1370520	A1	20031217	EP 2002-715150	20020319 <
R: AT, BE, CH,	DE,	DK, ES, FR,	GB, GR, IT, LI, LU,	NL, SE, MC, PT,
IE, SI, LT,	LV,	FI, RO, MK,	CY, AL, TR	
CN 1498205	A	20040519	CN 2002-806895	20020319 <
CN 100537527	С	20090909		
HU 2004000252	A2	20040830	HU 2004-252	20020319 <
JP 2004526736	Τ	20040902	JP 2002-576192	20020319 <
JP 4557492	В2	20101006		
NZ 527680	Α	20050729	NZ 2002-527680	20020319 <
ZA 2003006727	Α	20041129	ZA 2003-6727	20030828 <
MX 2003008484	A	20031208	MX 2003-8484	20030919 <
JP 2010001316	A	20100107	JP 2009-232046	20091005 <
PRIORITY APPLN. INFO.:			US 2001-277534P	P 20010321 <
			JP 2002-576192	A3 20020319 <
			WO 2002-US8300	W 20020319

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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 137:263068

GΙ

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Title compds. [I; A = (un)substituted aryl, pyridinyl, pyrazinyl, pyridazinyl; Z = biphenylcarbamoyl, biphenylcarbonyl, biphenylsulfonyl; M = H, Me, Et, iso-Pr, n-Pr, cyclobutyl; n = 2-4; p = 1-6; R1 = NH2, NHR, NR2, NOR2, NH(CH2)nNR2; R = Me, Et, n-Pr, iso-Pr, cyclobutyl; R2 = H, alkyl] are prepared as antagonists of the Melanin-concentrating hormone (MCH) receptor. In one embodiment, this invention provides methods of preparing title compds., pharmaceutical compns. containing one or more of title compds., methods of preparing pharmaceutical formulations comprising one or more title compds., and methods of treatment, prevention or amelioration or one or more of diseases associated with the MCH receptor. Thus, the title compound II was an illustrative inventive compound

IT 463936-60-9P 463936-71-2P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

 $\hbox{ (preparation of aryl and biaryl derivs. having Melanin-concentrating hormone }$

modulatory activity)

RN 463936-60-9 CAPLUS

CN Carbamic acid, (3,5-dichlorophenyl)-, 3-(dimethylamino)-1-[4-(4-pyridinyl)phenyl]propyl ester (9CI) (CA INDEX NAME)

RN 463936-71-2 CAPLUS

CN Carbamic acid, (3,5-dichlorophenyl)-, 3-(dimethylamino)-1-[4-(3-pyridinyl)phenyl]propyl ester (9CI) (CA INDEX NAME)

OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD

(4 CITINGS)

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 22 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2002:615623 CAPLUS Full-text

DOCUMENT NUMBER: 137:169517

TITLE: Oxazolyl-pyrazole derivatives as protein kinase

inhibitors, their preparation and combinatorial libraries, and their pharmaceutical use in the

treatment of cancer and other diseases and disorders

INVENTOR(S): Berta, Daniela; Felder, Eduard; Vulpetti, Anna; Villa,

Marzia

PATENT ASSIGNEE(S): Pharmacia Italia S.p.A., Italy

SOURCE: PCT Int. Appl., 107 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.					KIND DATE				APPL	ICAT	ION	DATE					
WO :	20020	16280) 4		A1	A1 20020815				 √O 20	 02-Е	P995			20	0201	 28 <
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FΙ,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	ΚP,	KR,	KΖ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TN,	TR,	TT,	TZ,
		UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZM,	ZW							
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AT,	BE,	CH,

CY,	DE, DK		•	, GR, IE, IT, LU, MC,		PT, SE, TR,
BF,	BJ, CF	CG,	CI, CM, G	, GN, GQ, GW, ML, MR,	ΝE,	SN, TD, TG
CA 2437260		A1	2002081	CA 2002-2437260		20020128 <
AU 20022460	76	A1	2002081	AU 2002-246076		20020128 <
AU 20022460	76	В2	2007063	4		
EP 1377589		A1	2004010	7 EP 2002-714136		20020128 <
EP 1377589		В1	200509	7		
R: AT,	BE, CH	DE,	DK, ES, FI	, GB, GR, IT, LI, LU,	NL,	SE, MC, PT,
IE,	SI, LT	LV,	FI, RO, M	, CY, AL, TR		
JP 200452039	94	T	2004070	3 JP 2002-563156		20020128 <
NZ 527123		A	2005042	NZ 2002-527123		20020128 <
AT 304017		T	2005091	AT 2002-714136		20020128 <
ES 2248532		Т3	2006031	ES 2002-714136		20020128 <
MX 200300686	63	A	2003111	MX 2003-6863		20030731 <
US 200401808	381	A1	2004091	US 2004-470859		20040415 <
US 7105535		В2	2006093	2		
PRIORITY APPLN. 1	INFO.:			GB 2001-2687	A	20010202 <
				WO 2002-EP995	W	20020128

<--

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 137:169517

GI

AB The method of treating protein kinase-linked diseases with oxazolyl-pyrazole derivs. I and their pharmaceutically acceptable salts is disclosed [wherein: R = H, alkyl, alkenyl, aryl, arylalkyl, (un)saturated cycloalkyl or cycloalkyloxy optionally condensed with 1 or 2 benzene rings, or optionally benzo-condensed 5- or 6-membered heterocyclyl or heterocyclylalkyl having 1 or 2 N/O/S atoms [all optionally substituted by one or more of: halo, NO2, cyano, OH, oxo, alkyl, alkoxyalkyl, perfluoroalkyl, (un)substituted aryl or

5- or 6-membered heterocyclyl having 1 or 2 N/O/S atoms, alkoxy, alkoxyalkyloxy, (un) substituted arylalkyloxy or aryloxy, alkylthio, alkylsulfonyl, arylthio, or arylsulfonyl, cycloalkyl, amino, alkylamino, dialkylamino, arylamino, alkylcarbonyl, alkyloxycarbonyl, alkylaminocarbonyl, aminocarbonyl, (un)substituted arylcarbonyl or heterocyclylcarbonyl, alkylcarbonylamino, alkyloxycarbonylamino, arylalkyloxycarbonylamino, arylcarbonylamino, aryloxycarbonylamino, carboxy, alkylcarbonyloxy, or arylcarbonyloxy]; Y = bond, CO, NHCO, SO2; WZ = benzo fusion, naphtho fusion, or an optionally benzocondensed 5- or 6-membered heterocycle having 1 or 2 N/O/S atoms, each optionally substituted by one or more of halo, nitro, cyano, alkyl, alkoxy, alkylsulfonyl, or aryl]. Also disclosed is a novel subset of I, including 382 individually named compds. I are useful in the treatment of diseases caused by and/or associated with an altered protein kinase activity, such as cancer, cell proliferative disorders, viral infections, autoimmune diseases and neurodegenerative disorders. Eleven examples are given, including solid-phase preparation of several compds. I and intermediates, and descriptions of 3 combinatorial libraries of 3874, 3172, and 2184 members, based on 4 claimed tables of reactants. For instance, Et 3-(3-nitrophenyl)pyrazole-4-carboxylate was bound to trityl chloride resin, saponified with NaOH in MeOH, and amidated with o-aminophenol. The resultant N-(2-hydroxyphenyl) amide was cyclized by Mitsunobu reaction to give a 1,3-benzoxazole derivative, followed by reduction of the nitro group to amino using SnCl2, amidation with PhCH2CO2H, and resin cleavage with TFA, to give title compound II. Inhibition assays against various kinases are described (no data).

N-[4-[4-(4-Methyl-7-isopropyl-1,3-benzoxazol-2-yl)pyrazol-3-yl]phenyl]-2-(2,3-dichlorophenoxy)acetamide 48187-45-92,

N-[3-[4-(Naphth[2,3-d]-1,3-oxazol-2-yl)pyrazol-3-yl]phenyl]-2-(2,3-dichlorophenoxy)acetamide

RL: CPN (Combinatorial preparation); PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); CMBI (Combinatorial study); PREP (Preparation); USES (Uses)

(drug candidate; preparation of oxazolylpyrazole derivs. as protein ${\tt kinase}$

inhibitors, and their combinatorial libraries and use as anticancer agents)

RN 448184-05-2 CAPLUS

CN Acetamide, 2-(2,3-dichlorophenoxy)-N-[4-[4-(6-methyl-2-benzoxazolyl)-1H-pyrazol-3-yl]phenyl]- (CA INDEX NAME)

RN 448185-96-4 CAPLUS

CN Acetamide, 2-(2,3-dichlorophenoxy)-N-[4-[4-[4-methyl-7-(1-methylethyl)-2-benzoxazolyl]-1H-pyrazol-3-yl]phenyl]- (CA INDEX NAME)

RN 448187-45-9 CAPLUS

CN Acetamide, 2-(2,3-dichlorophenoxy)-N-[3-(4-naphth[2,3-d]oxazol-2-yl-1H-pyrazol-3-yl)phenyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 14 THERE ARE 14 CAPLUS RECORDS THAT CITE THIS

RECORD (15 CITINGS)

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 23 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2002:555466 CAPLUS Full-text

DOCUMENT NUMBER: 137:125096

TITLE: Preparation of phenyl derivatives containing

inhibitors of coagulation factor for prophylaxis

and/or therapy of thromboembolic disorders

INVENTOR(S): Dorsch, Dieter; Mederski, Werner; Tsaklakidis,

Christos; Cezanne, Bertram; Gleitz, Johannes; Barnes,

Christopher

PATENT ASSIGNEE(S): Merck Patent G.m.b.H., Germany

SOURCE: PCT Int. Appl., 133 pp.

CODEN: PIXXD2

DOCUMENT TYPE: %%tent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	NO.			KIN	D	DATE			APPLICATION NO. DATE						APPLICATION NO.					
WO 2002	05723	 36		A1		20020	725	1	vo 20	01-E	P142	96		20	0112	05 <				
W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	ВG,	BR,	BY,	BZ,	CA,	CH,	CN,				
	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,				
	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KΕ,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,				
	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	ΝZ,	PH,	PL,				
	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,				
	US,	UZ,	VN,	YU,	ZA,	ZW														
RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AT,	BE,	CH,				
	CY,	DE,	DK,	ES,	FΙ,	FR,	GB,	GR,	ΙE,	ΙT,	LU,	MC,	NL,	PT,	SE,	TR,				
	BF,	ΒJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML_{\prime}	MR,	NE,	SN,	TD,	TG				
DE 1010																				
								(CA 20	01-2	4349	37		20011205 <-						
CA 2434	937			С		2010	1109													
AU 2002	22799	93		A1	2	20020	0730	Z	AU 20	02-2	2799	3		20	0112	05 <				
AU 2002	2279	93		В2		2007	0809													
EP 1351	938			A1	2	20031	1015	I	EP 20	01-9	8958	0		20	0112	05 <				
EP 1351	938			В1		2007	0411													
R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,				
	IE,	SI,	LT,	LV,	FΙ,	RO,	MK,	CY,	AL,	TR										
BR 2001																				
CN 1518	541			A	2	20040	0804 CN 2001-823061 20011205 <-							05 <						

JP	2004535362	T	20041125	JP 2002-557917		20011205 <
JP	4180375	B2	20081112			
HU	2005000110	A2	20050628	HU 2005-110		20011205 <
AT	359271	T	20070515	AT 2001-989580		20011205 <
ES	2284718	T 3	20071116	ES 2001-989580		20011205 <
MX	2003006483	A	20030922	MX 2003-6483		20030718 <
IN	2003KN01033	A	20060602	IN 2003-KN1033		20030813 <
ZA	2003006419	A	20041118	ZA 2003-6419		20030818 <
US	20040087582	A1	20040506	US 2003-466680		20031218 <
US	7273867	B2	20070925			
PRIORIT	Y APPLN. INFO.:			DE 2001-10102322	Α	20010119 <
				WO 2001-EP14296	W	20011205

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OTHER SOURCE(S): MARPAT 137:125096

AB Novel compds. of the formula R1R2C6H3-W-X-Y-T in which W, X, Y, T, R1 and R2 are as defined in Patent Claim 1, are inhibitors of coagulation factor Xa and can be employed for the prophylaxis and/or therapy of thromboembolic disorders. Thus, 3-(5-methyl-1,2,4-oxadiazol-3-yl)phenol wa reacted with Et 2-bromovalerate, sodium hydroxide, thionyl chloride,

4-morpholin-4-ylaniline, followed a hydrogenation in acetic acid to give 2-(3-amidinophenoxy)-N-(4-morpholin-4-ylphenyl) valeramide acetate, showing IC50=3x10-7 M and IC50=4.9x10-7 M.

IT 444002-21-5P 444002-22-6P

RL: IMF (Industrial manufacture); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of Ph derivs. containing inhibitors of coagulation factor

for

prophylaxis and/or therapy of thromboembolic disorders)

RN 444002-21-5 CAPLUS

CN Pentanamide,

N-[4-(2,5-dioxo-1-pyrrolidinyl)-3-(trifluoromethyl)phenyl]-2-[3-(5-methyl-1,2,4-oxadiazol-3-yl)phenoxy]- (CA INDEX NAME)

RN 444002-22-6 CAPLUS

CN Pentanamide, N-[3-chloro-4-(2,5-dioxo-1-pyrrolidinyl)phenyl]-2-[3-(5-methyl-1,2,4-oxadiazol-3-yl)phenoxy]- (CA INDEX NAME)

$$\begin{array}{c|c} Me & O & N & O & CH & C & NH & C1 & O \\ \hline O & N & O & CH & C & NH & C1 & O \\ \hline \end{array}$$

OS.CITING REF COUNT: 18 THERE ARE 18 CAPLUS RECORDS THAT CITE THIS

RECORD (18 CITINGS)

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 24 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2002:368463 CAPLUS Full-text

DOCUMENT NUMBER: 136:386109

TITLE: Preparation of amide derivatives as antiherpes agents

INVENTOR(S): Kontani, Toru; Miyata, Junji; Hamaguchi, Wataru;

Miyazaki, Yoji; Suzuki, Hiroshi; Nakai, Eiichi;

Kageyama, Shunji

PATENT ASSIGNEE(S): Yamanouchi Pharmaceutical Co., Ltd., Japan; Rational

Drug Design Laboratories

SOURCE: PCT Int. Appl., 71 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT		KIN	KIND DATE			APPLICATION NO.						DATE				
WO 2002	03855	 54		A1	_ 2	20020	 0516	V						20	0111	.08 <
W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	ВG,	BR,	BY,	BZ,	CA,	CH,	CN,
	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,
	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PH,	PL,
	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,
	US,	UZ,	VN,	YU,	ZA,	ZW										
RW:	GH,	GM,	ΚE,	LS,	MW,	MΖ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,
	DE,	DK,	ES,	FΙ,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,
	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG	
CA 2428	184			A1	2	20020)516		CA 20	01-2	4281	84		20	0111	.08 <
CA 2428	184			С		2010	0330									
AU 2002	01273	3 4		Α	2	20020)521	Z	AU 20	02-1	2734			20	0111	.08 <
EP 1340	750			A1	2	20030	0903	E	EP 20	01-9	8103	3		20	0111	.08 <
EP 1340	750			В1		2005	0817									
R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
	ΙE,	SI,	LT,	LV,	FΙ,	RO,	MK,	CY,	AL,	TR						
AT 3021	97			Τ	2	20050	915	Z	AT 20	01-9	8103	3		20	0111	.08 <
ES 2247	177			Т3	2	20060	0301	E	ES 20	01-9	8103	3		20	0111	.08 <
JP 3913	172			В2	2	20070	0509	Ċ	JP 20	02-5	4108	9		20	0111	.08 <
US 2004	00342	232		A1	2	20040	219	J	JS 20	03-4	1637	1		20	0305	12 <
US 6949	543			В2		2005	0927									
RIORITY APPLN. INFO.:					JP 2000-344354				А	A 20001110 <						
							,	WO 2	001-	JP97	90	Ţ	w 2	0011	108	
_																

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 136:386109

GΙ

$$\mathbb{R}^{1} \xrightarrow{\mathbb{N}} \mathbb{R}^{2}$$

AB The title compds. I [R1, R2 = H, alkyl, etc.; ring A = (un)substituted aryl, etc.; X = CO, SO2; R3 = (un)substituted cycloalkyl, etc.] are prepared These amide derivs. are useful as drugs and antiviral agents, in particular, preventives or remedies for various diseases caused by the infection with herpesviruses, more specifically, various herpesvirus infections such as pox (blister) caused by the infection with varicella zoster virus, herpes caused by the recurrent infection with latent varicella zoster virus, herpes labialis and herpes encephalitis caused by the infection with HSV-1 and genital herpes caused by the infection with HSV-2.

N-([[4-(2-Aminothiazol-4-yl)phenyl]carbamoyl]methyl)-4-fluoro-N-(2,3-dihydro-1H-indol-6-yl)benzamide dihydrochloride showed EC50 value of 0.046 μ M against varicella zoster virus, vs. EC50 value of 4.3 μ M shown by acyclovir.

IT 425688-62-69

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of amide derivs. as antiherpes agents)

RN 425688-62-6 CAPLUS

CN Benzamide, 4-fluoro-N-(4-fluorophenyl)-N-[2-oxo-2-[[4-[2-(2-pyridinylamino)-4-thiazolyl]phenyl]amino]ethyl]-, hydrochloride (1:2)

(CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

●2 HC1

OS.CITING REF COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (8 CITINGS)

REFERENCE COUNT: 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L87 ANSWER 25 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN
ACCESSION NUMBER:
                        2002:72061 CAPLUS Full-text
DOCUMENT NUMBER:
                        136:118465
TITLE:
                        Preparation of 2-aryldihydroxypyrimidine-4-carboxylic
                        acids as hepatitis C viral polymerase inhibitors
INVENTOR(S):
                        Gardelli, Cristina; Giuliano, Claudio; Harper, Steven;
                        Koch, Uwe; Narjes, Frank; Ontoria Ontoria, Jesus
                        Maria; Poma, Marco; Ponzi, Simona; Stansfield, Ian;
                        Summa, Vincenzo
PATENT ASSIGNEE(S):
                        Istituto di Ricerche di Biologia Molecolare P.
                        Angeletti S.p.A., Italy
SOURCE:
                        PCT Int. Appl., 162 pp.
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
                        English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                      KIND DATE APPLICATION NO.
                                                               DATE
    PATENT NO.
     _____
                      ----
                                         -----
                              20020124 WO 2001-EP7955
    WO 2002006246
                       A1
                                                               20010711 <--
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT,
            RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US,
            UZ, VN, YU, ZA, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
            DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
            BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
    CA 2418288
                              20020124 CA 2001-2418288
                        Α1
                                                                20010711 <--
    EP 1309566
                              20030514 EP 2001-951664
                                                               20010711 <--
                        A1
                               20091007
    EP 1309566
                        B1
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                       T
                             20040212
                                        JP 2002-512150
    JP 2004504304
                                                                20010711 <--
    AU 2001272530
                        В2
                              20060803
                                          AU 2001-272530
                                                                20010711 <--
    AT 444956
                        Τ
                              20091015
                                          AT 2001-951664
                                                                20010711 <--
    US 20040106627
                       A1
                              20040603
                                         US 2003-333431
                                                                20030709 <--
                        B2
                               20060815
    US 7091209
PRIORITY APPLN. INFO.:
                                          GB 2000-17676
                                                             A 20000719 <--
                                          WO 2001-EP7955
                                                             W 20010711
<--
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OTHER SOURCE(S):
                        MARPAT 136:118465
     RR1 (R1 = 4-carboxy-5,6-dihydroxy-2-pyrimidinyl)[I; R = (un)substituted
     (hetero)aryl] were prepared Thus, 2-(O2N)C6H4C(:NOH)NH2 (preparation
     given) N-was alkenylated by MeO2CC.tplbond.CCO2Me and the product cyclized
     to give, after reduction, N-acylation, and saponification, I [R =
     2-(2-C1C6H4CH2NHCONH)C6H4]. Data for biol. activity of I were given.
                 865876-12-6 865876-13-7
1102359-49-8 1102360-10-0
ΙT
     865876-11-5
    866052-40-6
    1102360-16-6 1102360-19-9 1102360-21-3
    1102363-65-4 1102363-69-8 1102363-70-1
```

1102363-71-2 1102363-72-3 1102363-75-6

1102363-77-8 1102363-82-5

RL: PRPH (Prophetic)

(Preparation of 2-aryldihydroxypyrimidine-4-carboxylic acids as

hepatitis C viral polymerase inhibitors)

RN 865876-11-5 CAPLUS

CN 4-Pyrimidinecarboxylic acid, 2-[2-[[[(2-

fluorophenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6oxo- (CA INDEX NAME)

RN 865876-12-6 CAPLUS

CN 4-Pyrimidinecarboxylic acid, 2-[2-[[[[(2-bromophenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6-oxo- (CA INDEX NAME)

RN 865876-13-7 CAPLUS

CN 4-Pyrimidinecarboxylic acid, 2-[2-[[[(2-chloro-6-

methylphenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6oxo- (CA INDEX NAME)

$$\begin{array}{c} \text{Me} \\ \text{NH} \\ \text{C} \\ \text{NH} \\ \text{CH}_2 \\ \text{C}_1 \\ \text{HO}_2 \\ \text{C}_1 \\ \text{HO}_2 \\ \text{C}_1 \\ \text{C}_1 \\ \text{C}_1 \\ \text{C}_2 \\ \text{C}_1 \\ \text{C}_2 \\ \text{C}_1 \\ \text{C}_2 \\ \text{C}_1 \\ \text{C}_1 \\ \text{C}_2 \\ \text{C}_2 \\ \text{C}_1 \\ \text{C}_2 \\ \text{C}_2 \\ \text{C}_3 \\ \text{C}_4 \\ \text{C}_4 \\ \text{C}_5 \\ \text{C}_5 \\ \text{C}_6 \\ \text{C}_6 \\ \text{C}_6 \\ \text{C}_7 \\ \text{C}_7$$

RN 866052-40-6 CAPLUS

CN 4-Pyrimidinecarboxylic acid, 2-[2-[[[[(2-chlorophenyl)methyl]amino]carbonyl]amino]-5-[(2-thienylcarbonyl)amino]phenyl]-1,6-dihydro-5-hydroxy-6-oxo-(CA INDEX NAME)

RN 1102359-49-8 CAPLUS

CN 4-Pyrimidinecarboxylic acid, 2-[3-[[[[(3-

fluorophenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6oxo- (CA INDEX NAME)

RN 1102360-10-0 CAPLUS

CN 4-Pyrimidinecarboxylic acid, 2-[3-[[[(2-

chlorophenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6oxo- (CA INDEX NAME)

RN 1102360-16-6 CAPLUS CN INDEX NAME NOT YET ASSIGNED

RN 1102360-19-9 CAPLUS

CN 4-Pyrimidinecarboxylic acid, 2-[3-[[[[(3-

chlorophenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6oxo- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & \\ & & \\ &$$

RN 1102360-21-3 CAPLUS

CN 4-Pyrimidinecarboxylic acid, 2-[3-[[[(3,4-

dichlorophenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6oxo- (CA INDEX NAME)

$$\begin{array}{c} \text{CO}_{2H} \\ \text{NH} \\ \text{NH} \\ \text{C-NH-CH}_{2} \end{array} \begin{array}{c} \text{C1} \\ \text{C2} \\ \text{C2} \\ \text{C3} \\ \text{C4} \\ \text{C4} \\ \text{C1} \\ \text{C2} \\ \text{C2} \\ \text{C3} \\ \text{C4} \\ \text{C4} \\ \text{C5} \\ \text{C6} \\ \text{C6$$

RN 1102363-65-4 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

RN 1102363-69-8 CAPLUS CN INDEX NAME NOT YET ASSIGNED

RN 1102363-70-1 CAPLUS CN INDEX NAME NOT YET ASSIGNED

RN 1102363-71-2 CAPLUS
CN INDEX NAME NOT YET ASSIGNED

RN 1102363-72-3 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

RN 1102363-75-6 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

$$\begin{array}{c} C1 \\ NH - C - NH - CH_2 \\ HO_2C - HO_0 \end{array}$$

RN 1102363-77-8 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

$$\begin{array}{c|c} & \text{Me} & \text{CI} \\ & \text{NH} - \text{CH}_2 & \text{CI} \\ & \text{HO}_2\text{C} & \text{NH} - \text{CH}_2 \\ & \text{HO}_2\text{C} & \text{NH} - \text{CH}_2 \\ & \text{HO}_2\text{C} & \text{NH} - \text{CH}_2 \\ & \text{NH} - \text{CH}_2 & \text{NH} - \text{CH}_2 \\ & \text$$

RN 1102363-82-5 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

IT 391680-76-5P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

 $\hbox{(preparation of 2-aryldihydroxypyrimidine-4-carboxylic acids as hepatitis C}$

viral polymerase inhibitors)

RN 391680-76-5 CAPLUS

CN 4-Pyrimidinecarboxylic acid, 2-[2-[[[[(2-

chlorophenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6oxo- (CA INDEX NAME)

IT 391680-87-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

 $\hbox{(preparation of 2-aryldihydroxypyrimidine-4-carboxylic acids as hepatitis C}$

viral polymerase inhibitors)

RN 391680-87-8 CAPLUS

CN 4-Pyrimidinecarboxylic acid, 2-[2-[[[[(2-

chlorophenyl)methyl]amino]carbonyl]amino]phenyl]-1,6-dihydro-5-hydroxy-6oxo-, methyl ester (CA INDEX NAME)

$$\begin{array}{c} C1 \\ NH-C-NH-CH2 \end{array}$$

OS.CITING REF COUNT: 23 THERE ARE 23 CAPLUS RECORDS THAT CITE THIS

RECORD (25 CITINGS)

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 26 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2001:851793 CAPLUS Full-text

DOCUMENT NUMBER: 136:5986

TITLE: Preparation of azole inhibitors of cytokine production

INVENTOR(S): Bamaung, Nwe Y.; Basha, Anwer; Djuric, Stevan W.; Gubbins, Earl J.; Luly, Jay R.; Tu, Noah P.; Madar,

David J.; Warrior, Usha; Wiedeman, Paul E.; Zhou, Xun;

Sciotti, Richard J.; Wagenaar, Frank L.

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 124 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 20010044445	A1	20011122	US 1999-289155	19990408 <
PRIORITY APPLN. INFO.:			US 1999-289155	19990408 <
ACCICMMENT HICTORY FOR	HC DATEN	T ATATIADIR	TH TOUC DIODIAY EODMAT	

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 136:5986

GI

$$\begin{array}{c|c}
R^2 & R^3 \\
R^4 & R^4 \\
N - Q - E \\
R^5 & R^5
\end{array}$$

$$_{\mathrm{F_{3}C}}$$
N NH-CO II

The title compds. [I; R1, R3 = H, aryl, perfluoroalkyl, etc.; Z = N, C; R2 is absent or = H, alkyl, cycloalkyl, etc.; Q = (hetero)aryl (when Q = Ph, the Ph is 2-, 3-, or 4-substituted by E relative to the position of attachment of the pyrazole or 1,2,4-triazole ring to the Ph ring); R4, R5 = H, alkyl, haloalkyl, etc.; E = NO2, NH2, etc.], useful for inhibiting cytokine (Interleukin-2, Interleukin-4, or Interleukin-5) production and cellular proliferation in stimulated human T cell lines or human peripheral blood mononuclear cells (biol. data given) and therefore have utility in the treatment of diseases that are prevented by or ameliorated with cytokine inhibitors, were prepared General procedures for preparation of compds. I were described. Thus, the title compound II was prepared

IT 245746-03-69

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of azole inhibitors of cytokine production)

RN 245746-03-6 CAPLUS

CN Propanamide, N-[4-[3,5-bis(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]-2-(4-chlorophenoxy)-2-methyl- (CA INDEX NAME)

$$F_3C \xrightarrow{N}_{CF_3} NH \xrightarrow{O Me}_{Me} C1$$

OS.CITING REF COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS)

L87 ANSWER 27 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2001:851123 CAPLUS Full-text

DOCUMENT NUMBER: 136:5985

TITLE: Preparation of tricyclic pyrazole derivatives as

tyrosine kinase inhibitors for treatment of

angiogenesis-related diseases

INVENTOR(S): Doyle, Kevin J.; Rafferty, Paul; Steele, Robert W.;

Wilkins, David J.; Arnold, Lee D.; Hockley, Michael; Ericsson, Anna M.; Iwasaki, Nobuhiko; Ogawa, Nobuo

PATENT ASSIGNEE(S): Knoll G.m.b.H., Germany SOURCE: PCT Int. Appl., 183 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

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	WO	2001	08784	16		A2				V					20010517 <				
		W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DΖ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MΖ,	NO,	NZ,	PL,	PT,	
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			UΖ,	VN,	YU,	ZA,	ZW												
		RW:	GH,	GM,	ΚE,	LS,	MW,	MΖ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,	
			DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,	
			ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	G₩,	ML,	MR,	ΝE,	SN,	TD,	ΤG			
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							WO 2001-US16153				1	W 20010517							
_																			

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 136:5985

GI

$$\begin{bmatrix} A \\ N \end{bmatrix} \begin{bmatrix} X \\ N \end{bmatrix} \begin{bmatrix} B \\ M \end{bmatrix} \begin{bmatrix} R1 \end{bmatrix} \begin{bmatrix} R1 \end{bmatrix}$$

AB Title compds. I [m = 1-10; X = (CH2)n, CO, O, C:NOR10, NR11, (CH2)n, S, SO, or SO2; n = 1-3; R10 = alkyl; R11 = (un)substituted alkyl or Ph; B = (cyclo)alkyl, aryl, pyridyl, thienyl, furyl, or pyrrolyl; R1 = H, halo, OH, NO2, CN, hydroxyamidino, CH2NH2, formamidomethyl, (un)substituted

alkenyl(oxy), alkynyl, or YW; Y = absent or alkyl, alkoxy, O, S, or CO; W = H, OH, (un)substituted Ph, alkoxy, or amino; ring A is optionally substituted with halo, OH, NO2, CN, or (un)substituted alkyl, alkoxy, PhO, carboxy, carbamoyl, amino, amido, aralkyl, alkenyl, or alkynyl; with provisos; and racemic mixts., racemic diastereomeric mixts., tautomers, optical isomers, and pharmaceutically acceptable salts thereof] were prepared as protein kinase inhibitors, especially tyrosine kinase inhibitors. Thus, indan-1-one hydrazone (preparation given) in THF at 0° was treated with BuLi and then with Me 3,4,5-trimethoxybenzoate to give 3-(3,4,5-trimethoxyphenyl)-1,4-dihydroindeno[1,2-c]pyrazole. Example compds. significantly inhibited KDR kinase at concns. of $\leq 50~\mu\text{M}$.

IT 268563-63-9P, N-(2,5-Difluorobenzyl)-N'-[4-(1,4-

dihydroindeno[1,2-c]pyrazol-3-yl)phenyl]urea

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

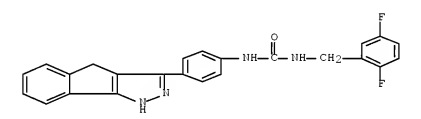
(preparation of tricyclic pyrazole derivs. as tyrosine kinase inhibitors

for

treatment of angiogenesis-related diseases)

RN 268563-63-9 CAPLUS

CN Urea, N-[(2,5-difluorophenyl)methyl]-N'-[4-(1,4-dihydroindeno[1,2-c]pyrazol-3-yl)phenyl]- (CA INDEX NAME)



OS.CITING REF COUNT: 8 THERE ARE 8 CAPLUS RECORDS THAT CITE THIS RECORD

(8 CITINGS)

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 28 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2001:833294 CAPLUS Full-text

DOCUMENT NUMBER: 135:357934

TITLE: Preparation of tetrazolylbenzoylureas as pesticides

and herbicides.

INVENTOR(S): Maurer, Fritz; Erdelen, Christoph PATENT ASSIGNEE(S): Bayer Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 85 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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20011115
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     WO 2001085705
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             HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
             LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO,
             RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ,
             VN. YU. ZA. ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
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PRIORITY APPLN. INFO.:
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                                             WO 2001-EP4899
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                                             IN 2001-MU402
                                                                 A3 20010503 <--
                                                                 A3 20021108 <--
                                             US 2002-275829
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 135:357934
GI

AB Title compds. [I; R1 = halo; R2 = H, halo; R3 = halo, alkyl, haloalkyl; n = 0-2; R4 = H, (substituted) alkyl, alkenyl, alkoxyalkyl, alkoxycarbonylalkyl, alkylsulfonyl, aryl, aralkyl, arylsulfonyl, cycloalkyl, cycloalkylalkyl, heterocyclyl, heterocyclylalkyl], were prepared Thus, 5-(3-chloro-4-aminophenyl)tetrazole (preparation given) in MeCN was treated with 2,6-difluorobenzoyl isocyanate in MeCN to precipitate 53% N-(2,6-difluorobenzoyl)-N'-(2-chloro-4-tetrazol-5-ylphenyl)urea. Several I at 0.1% on cabbage leaves gave 100% kill of Spodoptera frugiperda after 7 days.

1139482-71-5 ΙT 1139482-69-1 1139482-72-6 1139482-74-8 1139482-73-7 1139482-77-1 1139482-79-3 1139482-78-2 1139482-80-6 RL: PRPH (Prophetic) (Preparation of tetrazolylbenzoylureas as pesticides and herbicides.) RN 1139482-69-1 CAPLUS CN Benzamide, 2-chloro-N-[[[4-chloro-3-[1-(difluoromethyl)-1H-tetrazol-5yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 1139482-71-5 CAPLUS
CN Benzamide, N-[[[3-[1-(difluoromethyl)-1H-tetrazol-5-yl]-5(trifluoromethyl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 1139482-72-6 CAPLUS
CN Benzamide, 2-chloro-N-[[[3-[1-(difluoromethyl)-1H-tetrazol-5-yl]-5-(trifluoromethyl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 1139482-73-7 CAPLUS
CN Benzamide, N-[[[2-chloro-5-[1-(difluoromethyl)-1H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 1139482-74-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[2-chloro-5-[1-(difluoromethyl)-1H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 1139482-77-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-[1-(difluoromethyl)-1H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 1139482-78-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-(2H-tetrazol-5-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 1139482-79-3 CAPLUS

CN Benzamide, 2,6-difluoro-N-[[[3-(2H-tetrazol-5-yl)phenyl]amino]carbonyl]-(CA INDEX NAME)

RN 1139482-80-6 CAPLUS

CN Benzamide, N-[[[4-chloro-3-[1-(difluoromethyl)-1H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

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ΙT
     372192-41-12
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                                    372192-44-4P
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RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of tetrazolylbenzoylureas as pesticides and herbicides)

RN 372192-41-1 CAPLUS

CN Benzamide, N-[[[2-chloro-4-(2H-tetrazol-5-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & &$$

RN 372192-43-3 CAPLUS

CN Benzamide, N-[[[2-chloro-4-[2-(tetrahydro-2H-pyran-2-yl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-44-4 CAPLUS

CN Benzamide, N-[[[2-chloro-4-(1-methyl-1H-tetrazol-5-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 372192-46-6 CAPLUS

CN Benzamide, N-[[[2-chloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-47-7 CAPLUS

CN Benzamide, N-[[[2-chloro-4-(2-methyl-2H-tetrazol-5-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 372192-48-8 CAPLUS

CN Benzamide, N-[[[4-chloro-2-(2H-tetrazol-5-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-49-9 CAPLUS

CN Benzamide, N-[[[2,5-dichloro-4-(2H-tetrazol-5-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-50-2 CAPLUS

CN Benzamide, N-[[[2-chloro-4-[2-(1-methylcyclohexyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-51-3 CAPLUS

CN Benzamide, N-[[[2-chloro-4-[2-[(tetrahydro-2H-pyran-2-yl)methyl]-2H-

tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c}
F & O & O & C1 \\
\hline
V & NH & CH2 & D
\end{array}$$

RN 372192-52-4 CAPLUS

CN Benzamide, 2-chloro-N-[[[2-chloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-6-fluoro- (CA INDEX NAME)

RN 372192-53-5 CAPLUS

CN Benzamide, N-[[[4-chloro-2-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-54-6 CAPLUS

CN 2H-Tetrazole-2-acetic acid, 5-[3-chloro-4-[[[(2,6-difluorobenzoyl)amino]carbonyl]amino]phenyl]-, methyl ester (CA INDEX NAME)

$$MeO = C + 2$$

$$NH = C + NH = C$$

$$NH = C + NH = C$$

$$NH = C + NH = C$$

RN 372192-55-7 CAPLUS

CN Benzamide, N-[[[4-[2-[2-(acetyloxy)ethyl]-2H-tetrazol-5-yl]-2-chlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-56-8 CAPLUS

CN Benzamide, N-[[[2-chloro-4-[2-(ethoxymethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-57-9 CAPLUS

CN Benzamide, N-[[[2-chloro-4-(2-propyl-2H-tetrazol-5-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & &$$

RN 372192-58-0 CAPLUS

CN Benzamide, N-[[[2,5-dichloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-59-1 CAPLUS

CN Benzamide, N-[[[4-[2-[(4-bromophenyl)methyl]-2H-tetrazol-5-yl]-2-

chlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-60-4 CAPLUS

CN Benzamide, N-[[[2-chloro-4-[2-(difluoromethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-61-5 CAPLUS

CN Benzamide, N-[[[2-chloro-4-(2-phenyl-2H-tetrazol-5-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-62-6 CAPLUS

CN Benzamide, N-[[[2-chloro-4-[1-(difluoromethyl)-1H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-63-7 CAPLUS

CN Benzamide, N-[[[2-chloro-4-[2-(2-methylpropyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$i-Bu = N$$

$$N = N$$

RN 372192-64-8 CAPLUS

CN Benzamide,

N-[[[2-chloro-4-[2-(3,4,4-trifluoro-3-buten-1-yl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c} \text{CF2} \\ \text{F-C-CH2-CH2} \\ \end{array}$$

RN 372192-65-9 CAPLUS

CN Benzamide, N-[[[2,6-dichloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$t-Bu = N$$

$$N = N$$

$$C1$$

$$NH = C$$

RN 372192-66-0 CAPLUS

CN Benzamide, N-[[[2,6-dichloro-4-[2-(2-methylpropy1)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-67-1 CAPLUS

CN Benzamide, N-[[[2,5-dichloro-4-[2-(2-methylpropy1)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-68-2 CAPLUS

CN Benzamide, N-[[[2,5-dichloro-4-(2-propyl-2H-tetrazol-5-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-69-3 CAPLUS

CN Benzamide, N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-70-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-6-fluoro- (CA INDEX NAME)

RN 372192-71-7 CAPLUS

CN Benzamide, N-[[[2,5-dichloro-4-[2-(difluoromethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-72-8 CAPLUS

CN Benzamide, N-[[[2-chloro-4-[2-(2-methyl-2-propen-1-yl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c} \text{CH2} \\ \text{Me-C-CH2} \\ \end{array}$$

RN 372192-73-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372192-74-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[2-chloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$t^{-Bu} = N$$

$$N = N$$

$$N = N$$

$$N = N$$

$$C = NH = C$$

RN 372192-75-1 CAPLUS

CN Benzamide,

2-chloro-N-[[[2,6-dichloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$t-Bu = N$$

$$N = N$$

$$C1$$

$$C1$$

$$C1$$

$$C1$$

RN 372192-76-2 CAPLUS

CN Benzamide,

2-chloro-N-[[[2,5-dichloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372192-77-3 CAPLUS

CN Benzamide, N-[[[3-chloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-78-4 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-6-fluoro- (CA INDEX NAME)

RN 372192-79-5 CAPLUS

CN Benzamide, N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]-3-fluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-80-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]-3-fluorophenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372192-81-9 CAPLUS

CN Benzamide, N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]-2,6-difluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-82-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]-2,5-difluorophenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372192-83-1 CAPLUS

CN Benzamide, N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]-2,5-difluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-84-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]-2,6-difluorophenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372192-85-3 CAPLUS

CN Benzamide, N-[[[2,5-dichloro-4-[2-(1-ethyl-1-methylpropyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-86-4 CAPLUS

CN Benzamide, N-[[[4-[2-(1-ethyl-1-methylpropyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c} Me \\ Et - C \\ Et \end{array} \qquad \begin{array}{c} NH - C \\ NH - C \\ \end{array}$$

RN 372192-87-5 CAPLUS

CN Benzamide, N-[[[4-[2-(1,1-dimethylpropy1)-2H-tetrazol-5-y1]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-88-6 CAPLUS

CN Benzamide, N-[[[4-[2-(1-ethyl-1-methylpropyl)-2H-tetrazol-5-yl]-2-fluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} Me \\ Et - C \\ Et \end{array} \qquad \begin{array}{c} Me \\ NH - C \\$$

RN 372192-89-7 CAPLUS

CN Benzamide, N-[[[4-[2-(1,1-dimethylpropy1)-2H-tetrazol-5-y1]-2-fluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c} \text{Me} \\ \text{Et} \underbrace{\begin{array}{c} \text{Me} \\ \text{Me} \end{array}}_{\text{Me}} \text{N} \underbrace{\begin{array}{c} \text{N} \\ \text{N} \\ \text{N} \end{array}}_{\text{N}} \text{N} \underbrace{\begin{array}{c} \text{F} \\ \text{N} \\ \text{F} \end{array}}_{\text{N}} \text{N} \underbrace{\begin{array}{c} \text{F} \\ \text{N} \\ \text{N} \end{array}}_{\text{N}} \text{N} \underbrace{\begin{array}{c} \text{F} \\ \text{N} \end{array}}_{\text{N}} \text{N} \underbrace{\begin{array}{c} \text{F}$$

RN 372192-90-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(1,1-dimethylpropy1)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c} \text{Me} \\ \text{Et} - \overset{\text{N}}{\overset{\text{L}}{\overset{\text{N}}}{\overset{\text{N}}{\overset{\text{N}}{\overset{\text{N}}{\overset{\text{N}}{\overset{\text{N}}}{\overset{\text{N}}{\overset{\text{N}}}{\overset{\text{N}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}{\overset{\text{N}}}{\overset{\text{N}}{\overset{\text{N}}}{\overset{\text{N}}{\overset{\text{N}}}{\overset{\text{N}}{\overset{\text{N}}}{\overset{\text{N}}{\overset{\text{N}}}{\overset{\text{N}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}}{\overset{\text{N}}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{\text{N}}}{\overset{N}}{\overset{N}}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N}}{\overset{N$$

RN 372192-91-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(1-ethyl-1-methylpropyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$Et = \begin{bmatrix} Me & & & \\ Et & & & \\ Et & & & \\ \end{bmatrix}$$

RN 372192-92-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(1,1-dimethylpropyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-6-fluoro- (CA INDEX NAME)

$$\begin{array}{c|c} Me & & \\ Et & C & \\ Me & & \\ N = & N \end{array}$$

RN 372192-93-3 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(1-ethyl-1-methylpropyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-6-fluoro- (CA INDEX NAME)

$$Et = \begin{bmatrix} Me & & & \\ & \ddots & & \\ Et & & N = & N \end{bmatrix}$$

$$NH = \begin{bmatrix} 0 & & & \\ & & & \\ & & & \\ & & & \end{bmatrix}$$

RN 372192-94-4 CAPLUS

CN Benzamide, N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]-3,5-difluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-95-5 CAPLUS

CN Benzamide, N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]-2,3-difluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-96-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]-2,3-difluorophenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372192-97-7 CAPLUS

CN Benzamide, N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]-2-fluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372192-98-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]-2-fluorophenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372192-99-9 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$t-Bu = N + C + N + C$$

RN 372193-00-5 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372193-01-6 CAPLUS

CN Benzamide, N-[[[2,3-dichloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372193-02-7 CAPLUS

CN Benzamide,

2-chloro-N-[[[2,3-dichloro-4-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372193-03-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[2-chloro-4-[2-(difluoromethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372193-04-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[2-chloro-4-[1-(difluoromethyl)-1H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372193-05-0 CAPLUS

CN Benzamide, N-[[[4-[2-(difluoromethy1)-2H-tetrazo1-5-y1]pheny1]amino]carbony1]-2,6-difluoro- (CA INDEX NAME)

RN 372193-06-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[2-(difluoromethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372193-07-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[1-(difluoromethyl)-1H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372193-08-3 CAPLUS

CN Benzamide, N-[[[4-[1-(difluoromethyl)-1H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372193-09-4 CAPLUS

CN Benzamide,

2-chloro-N-[[[2,5-dichloro-4-[2-(difluoromethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F_{2}CH = N \\ N \\ N \\ N \\ C1 \\ C1 \\ C1$$

RN 372193-10-7 CAPLUS

CN Benzamide, N-[[[3-chloro-4-[2-(difluoromethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372193-11-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-[2-(difluoromethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372193-12-9 CAPLUS

CN Benzamide, N-[[[4-[2-(difluoromethyl)-2H-tetrazol-5-yl]-3-fluorophenyl]amino]carbonyl]-2,6-difluoro-(CA INDEX NAME)

RN 372193-13-0 CAPLUS

CN Benzamide, N-[[[3-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372193-14-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-[2-(1,1-dimethylethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 372193-15-2 CAPLUS

CN Benzamide, N-[[[3-[2-(difluoromethyl)-2H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 372193-16-3 CAPLUS

CN Benzamide, N-[[[3-[1-(difluoromethyl)-1H-tetrazol-5-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD

(5 CITINGS)

REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 29 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2001:730744 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 135:288790

TITLE: Pyrrolopyrimidines as tyrosine kinase inhibitors

INVENTOR(S): Hirst, Gavin C.; Calderwood, David; Munschauer,

Rainer; Arnold, Lee D.; Johnston, David N.; Rafferty,

Paul

PATENT ASSIGNEE(S): Basf Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 453 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.						KIND DATE			APPL	ICAT	ION	DATE							
WO 2001072751					A1 20011004			WO 2000-US8593					20000329 <							
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,			
															GM,					
		ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LU,			
		LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,			
		SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW		
	RW:	GH,	GM,	ΚE,	LS,	MW,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,	DE,			
		DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,			
		CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG							
PRIO	PRIORITY APPLN. INFO.:						WO 200							00-US8593 20000329 <						
OTHER SOURCE(S):					MAR:	MARPAT 135:288790														
GT																				

AΒ Chemical compds. having structural formula I and physiol. acceptable salts and metabolites thereof, are inhibitors of serine/threonine and tyrosine kinase activity. Several of the kinases, whose activity is inhibited by these chemical compds., are involved in immunol., hyperproliferative, or angiogenic processes. Thus, these chemical compds. can ameliorate disease states where angiogenesis or endothelial cell hyperproliferation is a factor. These compds. can be used to treat cancer and hyperproliferative disorders, rheumatoid arthritis, disorders of the immune system, transplant rejections and inflammatory disorders. All exemplified compds. significantly inhibited either FGFR, PDGFR, KDR, Tie-2, Lck, Fyn, Blk, Lyn, or Src at $\leq 50~\mu\text{M}$, and some significantly inhibited cdc2 at $\leq 50~\mu\text{M}$. In I, ring A is a six membered aromatic ring or a five or six membered heteroarom. ring which is optionally substituted. L is -0-, -S-, -S(0)-, -S(0)2-, -N(R)-, -N[C(O)OR]-, -N[C(O)R]-, -N(SO2R)-, -CH2O-; -CH2S-, -CH2N(R)-, -C(NR) - ; -CH2N[C(O)R] - , -CH2N[C(O)OR] - , -CH2N(SO2R) - , -CH(NHR) - ,-CH[NHC(0)R]-, -CH(NHSO2R)-, -CH[NHC(0)OR]-, -CH[OC(0)R]-, -CH[OC(0)NHR]-, -CH:CH-; -C(:NOR)-, -C(O)-, -CH(OR)-, -C(O)N(R)-, -N(R)C(O)-, -N(R)S(O)-,-N(R)S(O)2-, -OC(O)N(R)-, -N(R)C(O)N(R)-, -NRC(O)O-, -S(O)N(R)-, -S(0)2N(R)-, -N[C(0)R]S(0)-, -N[C(0)R]S(0)2-, -N(R)S(0)N(R)-, -N(R)S(O)2N(R)-, -C(O)N(R)C(O)-, -S(O)N(R)C(O)-, -S(O)2N(R)C(O)-, -OS(0)N(R) -, -OS(0)2N(R) -, -N(R)S(0)O -, -N(R)S(0)2O -, -N(R)S(0)C(0) -, -N(R)S(O)2C(O)-, -SON[C(O)R]-, -SO2N[C(O)R]-, -N(R)SON(R)-, -N(R)SO2N(R)-, -C(0)O-, -N(R)P(OR')O-, -N(R)P(OR')-, -N(R)P(O)(OR')O-, -N(R)P(O)(OR')-, -N(R)P(O)(OR')-, -N(R)P(O)(OR')O-, -N-N[C(O)R]P(OR')O-, -N[C(O)R]P(OR')-, -N[C(O)R]P(O)(OR')O-, -N[C(O)R]P(OR')-, -CH(R)S(O)-, or -CH(R)S(O)2-. L is also -CH(R)N[C(O)OR]-, -CH(R)N[C(O)R]-, -CH(R)N(SO2R), -CH(R)O-, -CH(R)S-, -CH(R)N(R) -, -CH(R)N[C(O)R] -, -CH(R)N[C(O)OR] -, -CH(R)N(SO2R) -, -CH(R)C(:NOR)-, -CH(R)C(O)-, -CH(R)CH(OR)-, -CH(R)C(O)N(R)-,

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-CH(R)N(R)C(O) -, -CH(R)N(R)S(O) -, -CH(R)N(R)S(O)2 -, -CH(R)OC(O)N(R) -,
-CH(R)N(R)C(O)N(R) -, -CH(R)N(R)C(O)O -, -CH(R)S(O)N(R) -, -CH(R)S(O)2N(R) -,
-\mathrm{CH}(R)\,\mathrm{N}\,[\mathrm{C}(O)\,R]\,\mathrm{S}(O)\,-\,,\quad -\mathrm{CH}\,(R)\,\mathrm{N}\,[\mathrm{C}(O)\,R]\,\mathrm{S}(O)\,2-\,,\quad -\mathrm{CH}\,(R)\,\mathrm{N}\,(R)\,\mathrm{S}(O)\,\mathrm{N}\,(R)\,-\,,
-CH(R)N(R)S(O)2N(R)-, -CH(R)C(O)N(R)C(O)-, -CH(R)S(O)N(R)C(O)-,
-CH(R)S(O)2N(R)C(O)-, -CH(R)OS(O)N(R)-, -CH(R)OS(O)2N(R)-,
-CH(R)N(R)S(O)O-, -CH(R)N(R)S(O)2O-, -CH(R)N(R)S(O)C(O)-,
-CH(R)N(R)S(O)2C(O)-, -CH(R)SON[C(O)R]-, -CH(R)S(O)2N[C(O)R]-,
-CH(R)N(R)SON(R) -, -CH(R)N(R)S(O)2N(R) -; -CH(R)C(O)O -,
-CH(R)N(R)P(OR')O-, -CH(R)N(R)P(OR')-, -CH(R)N(R)P(O)(OR')O-,
-CH(R)N(R)P(O)(OR')-, -CH(R)N[C(O)R]P(OR')O-, -CH(R)N[C(O)R]P(OR')-,
-CH(R)N[C(0)R]P(0)(OR')O- or -CH(R)N[C(0)R]P(OR')-. In L, each R and R' is,
independently, -H, acyl, substituted or unsubstituted aliphatic, aromatic,
arylalkyl, heteroarom., cycloalkyl or arylalkyl; or L is -RbN(R)S(O)2-,
-RbN(R)P(O)-, or -RbN(R)P(O)O-, wherein Rb is an alkylene group which when
taken together with the sulfonamide, phosphinamide, or phosphonamide group
to which it is bound forms a five or six membered ring fused to ring A; or
L is II (X = O or nil; Y = O or nil) or III (Y = O, nil) wherein R85 taken
together with the phosphinamide, or phosphonamide is a 5-, 6-, or 7-membered,
aromatic, heteroarom. or heterocycloalkyl ring system. G is a direct bond,
-(CH2)j-(j = 1-6), C2-C6-alkenylene, C3-C8-cycloalkylene or
C1-C6-oxaalkylene group. R1 is substituted or optionally substituted
aliphatic, cycloalkyl, bicycloalkyl, cycloalkenyl, aromatic, heteroarom.,
heteroaralkyl, heterocycloalkyl, heterobicycloalkyl, alkylamido,
arylamido, -S(O)2-alkyl, -S(O)2-cycloalkyl, -C(O)alkyl, or -B-E, wherein B
is substituted or unsubstituted cycloalkyl, heterocycloalkyl, aromatic,
heteroarom., alkylene, aminoalkyl, alkylenecarbonyl, or aminoalkylcarbonyl
and E is substituted or unsubstituted azacycloalkyl, azacycloalkylcarbonyl,
azacycloalkylsulfonyl, azacycloalkylalkyl, heteroaryl,
heteroarylcarbonyl, heteroarylsulfonyl, heteroaralkyl, alkyl sulfonamido,
aryl sulfonamido, bicycloalkyl, ureido, thioureido or aryl. R2 is -H or
substituted or unsubstituted aliphatic, cycloalkyl, halogen, -OH, cyano,
aromatic, heteroarom., heterocycloalkyl, aralkyl, heteroaralkyl,
-(CH2)0-3NR4R5, or -(CH2)0-3C(O)NR4R5. R3 is substituted or unsubstituted
aliphatic, alkenyl, cycloalkyl, aromatic, heteroarom., or heterocycloalkyl
with provisos. R4, R5 and the N atom together form a 3, 4, 5, 6 or 7-membered,
substituted or unsubstituted heterocycloalkyl, heterobicycloalkyl or
heteroarom.; or R4 and R5 are each, independently, -H, azabicycloalkyl,
heterocycloalkyl, substituted or unsubstituted alkyl or Y-Z; Y is -C(0)-,
-(CH2)p-, -S(0)2-, -C(0)0-, -S02NH-, -C0NH-, -(CH2)p0-, -(CH2)pNH-,
-(CH2)pS-, -(CH2)pS(0)-, and -(CH2)pS(0)2-; p = 0-6; and Z is -H, or
substituted or unsubstituted alkyl, amino, arvl, heteroarvl or
heterocycloalkyl.
                    546 Example prepns. are included. For example, addition
of piperidine to
4-[4-amino-5-(4-phenoxyphenyl)-7H-pyrrolo[2,3-d]pyrimidin-7-
yl]cyclohexanone in DCE and AcOH, followed by treatment with Na[(AcO)3BH],
workup and chromatog., gave cis- and trans-IV.
262442-33-1P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); USES (Uses)
   (target compound; preparation of pyrrolopyrimidinamines as protein kinase
   inhibitors)
262442-33-1 CAPLUS
Acetamide, N-[4-[4-amino-7-[cis-4-(4-methyl-1-piperazinyl)cyclohexyl]-7H-
```

pyrrolo[2,3-d]pyrimidin-5-yl]-2-methoxyphenyl]-2-(4-chlorophenoxy)- (CA

ΙT

RN CN INDEX NAME)

Relative stereochemistry.

PAGE 1-A

PAGE 2-A

OS.CITING REF COUNT: THERE ARE 14 CAPLUS RECORDS THAT CITE THIS 14

RECORD (14 CITINGS)

THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 14 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 30 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2001:395257 CAPLUS Full-text

DOCUMENT NUMBER: 135:152703

TITLE: A convenient method for the preparation of

3-phenoxy/thiophenoxy-2(1H)quinolinones

AUTHOR(S): Hashim, S. Riaz; Reddy, P. Tirupathy

CORPORATE SOURCE: Organic Chemistry Division-I, Indian Institute of Chemical Technology, Hyderabad, 500 007, India

Indian Journal of Chemistry, Section B: Organic

Chemistry Including Medicinal Chemistry (2001),

40B(5), 357-360

CODEN: IJSBDB; ISSN: 0376-4699

National Institute of Science Communication, CSIR PUBLISHER:

DOCUMENT TYPE: Journal

SOURCE:

LANGUAGE: English

OTHER SOURCE(S): CASREACT 135:152703

AB A number of new 4-methyl-3-phenoxy(or thiophenoxy)-2(1H)quinolinones have been synthesized in excellent yields through a simple and efficient procedure involving reaction of N-(2-acetylphenyl)chloroacetamide with phenols and thiophenols and cyclization of

N-(2-acetylphenyl) phenoxy/thiophenoxyacetamide intermediates under basic conditions.

IT 352353-28-7P

RL: BYP (Byproduct); PREP (Preparation)
 (preparation of (thio)phenoxyquinolinones)

RN 352353-28-7 CAPLUS

CN Acetamide, 2-(2,6-dichlorophenoxy)-N-[2-(4-methyl-2-quinolinyl)phenyl]-(CA INDEX NAME)

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 31 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2001:300689 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 134:311208

TITLE: Preparation of pyrazolylphenylureas as 5-HT2A receptor

ligands

INVENTOR(S): Behan, Dominic P.; Beeley, Nigel R. A.; Chalmers,

Derek T.; Foster, Richard J.; Glen, Robert C.; Lawless, Michael S.; Liaw, Chen W.; Liu, Quin;

Menzaghi, Frederique; Russo, Joseph F.; Smith, Julian

R.; Thomsen, William J.

PATENT ASSIGNEE(S): Arena Pharmaceuticals, Inc., USA; Tripos, Inc.; et al.

SOURCE: PCT Int. Appl., 95 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 17

PATENT INFORMATION:

PATENT NO.				KIND DATE				APPLICATION NO.							DATE			
WO 2001029008			A1	20010426			WO 2000-US28347						20001013 <					
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PRIORITY APPLN. INFO.:
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                                                             A2 19990414 <--
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                                          US 1999-152708P
                                          WO 2000-US28347 W 20001013
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                                          AU 2002-219890
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                                          AU 2004-202147
                                                             A3 20040512
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                                          AU 2004-203102
                                                              A3 20040708
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OTHER SOURCE(S): MARPAT 134:311208
```

GΙ

Title compds. [I; R = C6H4[NR1C(:X)R9]-3; X = O, S, NR2; R9 = NR3R4, (CH2)mR5, O(CH2)nR6; R1-R3, R7 = H, alk(en)yl, cycloalkyl, etc.; R4-R6 = (cyclo)alkyl, aryl(methyl), etc.; R8 = halo, alk(en)yl, cycloalkyl, etc.; m, n = 0-4] were prepared Thus, I [R = C6H4(NHR10)-3, R7 = Me, R8 = Br] (II; R10 = H) was

amidated by 4-(MeS)C6H4NCS to give II [R10 = CSNHC6H4(SMe)-4]. Data for biol. activity of I were given.

IT 247038-30-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of pyrazolylphenylureas as 5-HT2A receptor ligands)

RN 247038-30-8 CAPLUS

CN Urea, N-[3-(4-bromo-1-methyl-1H-pyrazol-3-yl)phenyl]-N'-[(4-fluorophenyl)methyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 10 THERE ARE 10 CAPLUS RECORDS THAT CITE THIS

RECORD (10 CITINGS)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 32 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2001:185739 CAPLUS Full-text

DOCUMENT NUMBER: 134:237301

TITLE: Preparation of benzophenones and phenyl heteroaryl

ketones as inhibitors of reverse transcriptase

INVENTOR(S): Andrews, Clarence Webster; Chan, Joseph Howing;

Freeman, George Andrew; Romines, Karen Rene; Tidwell,

Jeffrey H.

PATENT ASSIGNEE(S): Glaxo Group Limited, UK; Pianetti, Pascal Maurice

Charles

SOURCE: PCT Int. Appl., 436 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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WO 20010)1798			A1 20010315			V	 √O 20		P84 8		20000831 <					
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JP 3739704	ь В2		3P 2001-321729	20000831 <				
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AU 770302	B2	20040130	AU 2000-77743	20000831 <				
CN 1636984	A	20050713	CN 2004-10095621	20000831 <				
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CN 1030300	C	20050713	CN 2000-815249	20000831 <				
AT 325106	T	20060615	AT 2000-967637	20000831 <				
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IE, SI, FI,		DI(, 15, 11)	GB, GR, 11, H1, H0,	NE, 52, 113, 11,				
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ZA 2002001664	A	20030527	ZA 2002-1664	20020227 <				
NO 2002001042	A	20020430	NO 2002-1042	20020301 <				
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MX 2002002347	A	20020812	MX 2002-2347	20020304 <				
US 7273863	В1	20070925	US 2002-70084	20020307 <				
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KR 2007093152	A	20070917	KR 2007-7019984	20070831 <				
PRIORITY APPLN. INFO.:			GB 1999-20872	A 19990904 <				
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ASSIGNMENT HISTORY FOR U			TAMAC					

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMATOTHER SOURCE(S):

MARPAT 134:237301

GI

$$\begin{array}{c|c}
 & R^2 & R^3 \\
 & N & R^4 \\
 & R^5 & R^3
\end{array}$$

II

The title compds. [I; X = C, O, N; R1 = alkyl, cycloalkyl, (un)substituted aryl, etc.; R2 = H, halo, alkyl; R3, R4 = H, OH, (un)substituted heterocyclyl, etc.; R5 = H, halo, alkyl, etc.], useful in the treatment of HIV infections, were prepared E.g., a 4-step synthesis of the ketone II which showed IC50 of between 101 nM and 1,000 nM against HIV-1 in MT4 cell assay, was described.

IT 329936-79-09 329936-85-89

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of benzophenones and Ph heteroaryl ketones as inhibitors of reverse transcriptase)

RN 329936-79-0 CAPLUS

CN Acetamide, 2-(2-benzoyl-4-chlorophenoxy)-N-[4-(1H-1,2,4-triazol-1-yl)phenyl]- (CA INDEX NAME)

RN 329936-85-8 CAPLUS

CN Acetamide, 2-(2-benzoyl-4-chlorophenoxy)-N-[4-(1,2,3-thiadiazol-4-yl)phenyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 18 THERE ARE 18 CAPLUS RECORDS THAT CITE THIS

RECORD (18 CITINGS)

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 33 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 2000:384159 CAPLUS Full-text

DOCUMENT NUMBER: 133:30670

TITLE: Preparation of substituted

benzo[de]isoquinoline-1,3-diones as glycoprotein IbIX

antagonists

INVENTOR(S): Mederski, Werner; Devant, Ralf; Barnickel, Gerhard;

Bernotat-Danielowski, Sabine; Melzer, Guido; Raddatz, Peter; Wu, Zhengdong; Dhanoa, Daljit; Soll, Richard;

Rinker, James; Graybill, Todd

PATENT ASSIGNEE(S): Merck Patent G.m.b.H., Germany

SOURCE: PCT Int. Appl., 278 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GI

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AΒ
     The title compds. [I; R = H, NO2; R1 = Het, -HetSO2Ar, NO2, etc.; R2 = Ar,
     Het1, -Het1Ar, etc.; Ar = Ph, biphenyl, pyridyl, etc.; Het, Het1 =
     (un) substituted (un) saturated mono-, bi- or tricyclic 5-13 membered
     heterocyclyl], useful as glycoprotein IbIX antagonists (no data) for the
     control of thrombotic disorders, were prepared and formulated. E.g.,
     preparation of II was given. Compds. I are effective at 0.02-10 mg/kg/day. 1098872-73-1 1098872-76-4 1098872-84-4
ΙT
     1098872-73-1
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        (Preparation of substituted benzo[de]isoquinoline-1,3-diones as
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RN
     1098872-73-1 CAPLUS
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RN 1098872-76-4 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

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RN 1098872-84-4 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

RN 1098872-88-8 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

$$H_2N-C-NH-CH_2$$

RN 1098872-89-9 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

$$H_2N-C-NH-CH_2$$

RN 1098872-90-2 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

$$H_2N-C-NH-CH_2$$

RN 1098873-22-3 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

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RN 1098873-23-4 CAPLUS
CN INDEX NAME NOT YET ASSIGNED

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RN 1098873-24-5 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

PAGE 1-A

PAGE 2-A

RN 1098873-29-0 CAPLUS

CN INDEX NAME NOT YET ASSIGNED

PAGE 1-A

PAGE 2-A

NH CH2

RN 1098873-33-6 CAPLUS CN INDEX NAME NOT YET ASSIGNED

PAGE 1-A

PAGE 2-A

RN 1098873-40-5 CAPLUS CN INDEX NAME NOT YET ASSIGNED

PAGE 2-A

RN 1098873-44-9 CAPLUS CN INDEX NAME NOT YET ASSIGNED

PAGE 2-A

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PAGE 2-A

RN 1098873-68-7 CAPLUS
CN INDEX NAME NOT YET ASSIGNED

PAGE 2-A

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CN INDEX NAME NOT YET ASSIGNED

RN 1098874-28-2 CAPLUS
CN INDEX NAME NOT YET ASSIGNED

PAGE 2-A

RN 1098874-41-9 CAPLUS CN INDEX NAME NOT YET ASSIGNED

PAGE 2-A

RN 1098876-45-9 CAPLUS CN INDEX NAME NOT YET ASSIGNED

PAGE 2-A

RN 1098876-48-2 CAPLUS CN INDEX NAME NOT YET ASSIGNED

RN 1098876-57-3 CAPLUS
CN INDEX NAME NOT YET ASSIGNED

RN 1098877-65-6 CAPLUS
CN INDEX NAME NOT YET ASSIGNED

PAGE 2-A

RN 1098879-51-6 CAPLUS CN INDEX NAME NOT YET ASSIGNED

RN 1098879-52-7 CAPLUS
CN INDEX NAME NOT YET ASSIGNED

RN 1098880-47-7 CAPLUS
CN INDEX NAME NOT YET ASSIGNED

PAGE 2-A

RN 1098880-51-3 CAPLUS
CN INDEX NAME NOT YET ASSIGNED

PAGE 2-A

RN 1098880-53-5 CAPLUS
CN INDEX NAME NOT YET ASSIGNED

RN 1098880-57-9 CAPLUS CN INDEX NAME NOT YET ASSIGNED

OS.CITING REF COUNT: 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD

(4 CITINGS)

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 34 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2000:335390 CAPLUS Full-text

DOCUMENT NUMBER: 132:347566

TITLE: Preparation of tricyclic pyrazole derivatives as

protein kinase inhibitors.

INVENTOR(S): Doyle, Kevin J.; Rafferty, Paul; Steele, Robert W.;

Wilkins, David J.; Hockley, Michael; Arnold, Lee D.;

Ericsson, Anna M.

PATENT ASSIGNEE(S): Basf Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 210 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

	TENT :							APPLICATION NO.					DATE				
WO	20000	02782	22		A2 20000518 A3 20000810			WO 1999-US26105						19991104 <			
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_	CA 2350235													19991104 <			
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EP	EP 1127051																
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_																	

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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 132:347566

GΙ

$$R^4$$
 R^5
 R^6
 R^1
 R^2

AB A method of inhibiting protein kinase activity comprises administration of title compds. [I; X = substituted methylene, CO, O, C:NOR7, NR8, (CH2)n, S,

SO, SO2; n = 1-3; R1 = H; R2 = (substituted) aryl, pyridyl, thienyl, furyl, pyrrolyl; R3-R6 = H, OH, halo, CO2H, alkoxycarbonyl, (substituted) alkyl, alkoxy, PhO, etc.; R7 = H, alkyl; with provisos]. Thus, indan-1-one hydrazone (preparation given) in THF at 0° was treated with BuLi and then with Me 3,4,5-trimethoxybenzoate to give 3-(3,4,5-trimethoxyphenyl)-1,4-dihydroindeno[1,2-c]pyrazole.

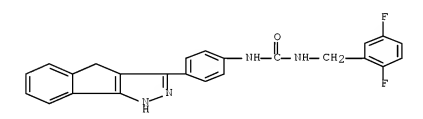
ΤТ 268563-63-9P

> RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

> (preparation of tricyclic pyrazole derivs. as protein kinase inhibitors)

RN 268563-63-9 CAPLUS

CN c]pyrazol-3-yl)phenyl]- (CA INDEX NAME)



OS.CITING REF COUNT: 23 THERE ARE 23 CAPLUS RECORDS THAT CITE THIS

RECORD (31 CITINGS)

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 35 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN 2000:247417 CAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 132:265193

TITLE: Preparation of phenylpyrazoles and hypolipidemic

agents

INVENTOR(S): Yamada, Hiroichi; Mochizuki, Nobuo; Uchida, Seiichi;

Umeda, Nobihiro

PATENT ASSIGNEE(S): Nippon Soda Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 19 pp. SOURCE:

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
JP 2000109465	А	20000418	JP 1999-221791		19990804 <
PRIORITY APPLN. INFO.:			JP 1998-222159	A	19980805 <
OTHER SOURCE(S):	CASRE	ACT 132:2651	93; MARPAT 132:2651	93	
GI					

$$\begin{array}{c}
N \\
N \\
N \\
R \\
12
\end{array}$$

$$\begin{array}{c}
N \\
R \\
1
\end{array}$$

$$N = NH$$

$$\downarrow_{\text{R12}} NH$$

$$\downarrow_{\text{R12}} NH$$

$$\downarrow_{\text{R12}} II$$

Title compds. I [R1 = H, C1-6 alkyl; X = C0, S02; A = (CR3R2)p(CR4:CR5)q; B = (CR6R7)r; R2, R3, R6, R7 = H, cyano, OH, halo, C1-6 alkyl, C1-6 alkoxy etc.; R4, R5 = H, C1-6 alkyl, C1-6 haloalkyl, (un)substituted benzyl; p, r = 0-6; q = 0-1; Y = O, S, SO, SO2, CO, etc.; n = 0-1; D = (un)substituted Ph; naphthyl, tetrahydronaphthyl, indanyl; R11 = halo, C1-6 alkyl, C1-6 alkoxy; m = 0-2; R12 = H, C1-6 alkyl] or their pharmaceutically acceptable salts are prepared by dehydration of pyrazoles II (R1, R11, R12, m = same as I) with HO2CAY1BD (A, B, Y, D, n = same as I). 5-(4-Aminophenyl)pyrazole (1.59 g) was reacted with 3.09 g benzoyl chloride in the presence of NEt3 in DMF at room temperature for 20 h to give 1.31 g phenyl-N-[4-(pyrazol-5-yl)phenyl]carboxamide showing in vivo good hypolipidemic activity.

IT 263257-79-0P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); IMF (Industrial manufacture); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

 $(\hbox{preparation of phenylpyrazoles by dehydration of aminophenylpyrazoles}$

and

carboxylic acids)

RN 263257-79-0 CAPLUS

CN Propanamide, 2-(4-chlorophenoxy)-2-methyl-N-[4-(1H-pyrazol-3-yl)phenyl]-(CA INDEX NAME)

$$\begin{array}{c|c} & \text{NH} & \text{O} & \text{Me} \\ & \text{NH} & \text{C} & \text{O} & \text{Me} \\ & \text{Me} & \text{O} & \text{Me} \\ \end{array}$$

L87 ANSWER 36 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 2000:210172 CAPLUS Full-text

DOCUMENT NUMBER: 132:251160

TITLE: Preparation of pyrrolopyrimidines as protein kinase

inhibitors

INVENTOR(S): Hirst, Gavin C.; Calderwood, David; Wishart, Neil;

Ritter, Kurt; Arnold, Lee D.

PATENT ASSIGNEE(S): Basf A.-G., Germany SOURCE: PCT Int. Appl., 304 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.				APPLICATION NO.	DATE		
W: AE, CZ, IN, MG, SL, RW: GH, DK,	03 AL, AM, DE, DK, IS, JP, MK, MN, TJ, TM, GM, KE, ES, FI, CI, CM,	AT, DM, KE, MW, TR, LS, FR, GA,	20000330 AU, AZ, BA, EE, ES, FI, KG, KP, KR, MX, NO, NZ, TT, TZ, UA, MW, SD, SL, GB, GR, IE, GN, GW, ML,	WO 1999-US21560 BB, BG, BR, BY, CA, GB, GD, GE, GH, GM, KZ, LC, LK, LR, LS, PL, PT, RO, RU, SD, UG, US, UZ, VN, YU, SZ, TZ, UG, ZW, AT, IT, LU, MC, NL, PT, MR, NE, SN, TD, TG CA 1999-2344249	CH, CN, CR, CU, HR, HU, ID, IL, LT, LU, LV, MD, SE, SG, SI, SK, ZA, ZW BE, CH, CY, DE, SE, BF, BJ, CF,		
AU 9960484		AI	20000330	AU 1999-60484	19990917 <		
AU 753555		В2	20021024	1333031			
EP 1114053		A1	20010711	19990917 <			
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			20020629 20040728	HU 2002-403	19990917 <		
HU 20020004 JP 20025265	.03	A3	20040728	ID 2000 E74112	10000017 /		
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US 20030153	750	A.	20030829	US 2000-537167	19990917 < 20000329 <		
US 6713474	752	A1 B2	20030614	05 2000-55/16/	20000329 <		
BG 105346		A	20011231	BG 2001-105346	20010315 <		
NO 20010013	56	A	20010516	NO 2001-1356	20010316 <		
ZA 20010022				ZA 2001-2204	20010316 <		
IN 2001CN00	376	A	20050304	IN 2001-CN376	20010316 <		
PRIORITY APPLN.	INFO.:			US 1998-100832P	P 19980918 <		
				US 1998-100833P	P 19980918		
<				US 1998-100834P	P 19980918		
<				US 1998-100946P	P 19980918		
<				WO 1999-US21560	W 19990917		
<							
ASSIGNMENT HISTO	RY FOR U	S PA	TENT AVAILAB	LE IN LSUS DISPLAY FO	ORMA'I'		

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 132:251160

GI

AΒ 7H-Pyrrolo[2,3-d]pyrimidin-4-amines (I) [wherein A = (un) substituted 6-membered aromatic ring or 5- or 6-membered heteroarom. ring; L = RbN(R)S(0)2, RbN(R)P(0), or RbN(R)P(0)0, where Rb = alkylene group which when taken together with the sulfonamide, phosphinamide or phosphonamide group to which it is bound forms a 5- or 6-membered ring fused to ring A, or L =5-, 6-, or 7-membered (oxa)azaphosphaarom. or (oxa)azaphosphacycloalkyl ring; R = H, acyl, or (un) substituted aliphatic, (hetero) aromatic, or cycloalkyl; R1 = (un)substituted (hetero)cyclic, (hetero)aromatic, amido, acyl, or (cyclo)alkylsulfonyl; R2 = H, halo, OH, CN, (un)substituted aliphatic, cycloalkyl, (hetero)aromatic, (hetero)aralkyl, amino, or amido; R3 (un) substituted aliphatic, alkenyl, (hetero) cycloalkyl, or (hetero)aromatic; n = 0-6], and physiol. acceptable salts and metabolites thereof, were prepared For example, addition of piperidine to 4-[4-amino-5-(4-phenoxyphenyl)-7H-pyrrolo[2,3-d]pyrimidin-7yl]cyclohexanone in DCE and AcOH, followed by workup and chromatog., gave cis- and trans-II. I inhibit serine/threonine and tyrosine kinase activity, which are involved in immunol., hyperproliferative, and angiogenic processes. All exemplified compds. significantly inhibited either FGFR, PDGFR, KDR, Tie-2, Lck, Fyn, Blk, Lyn, or Src at concns. of $\leq 50 \, \mu\text{M}$, and some significantly inhibited cdc2 at concns. of $50 \le \mu M$. Thus, these compds. are useful in the treatment of cancer and hyperproliferative disorders, rheumatoid arthritis, disorders of the immune system, transplant rejections, and inflammatory disorders.

IT 262442-33-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(target compound; preparation of 7H-pyrrolo[2,3-d]pyrimidin-4-amines as protein kinase inhibitors)

RN 262442-33-1 CAPLUS

CN Acetamide, N-[4-[4-amino-7-[cis-4-(4-methyl-1-piperazinyl)cyclohexyl]-7H-pyrrolo[2,3-d]pyrimidin-5-yl]-2-methoxyphenyl]-2-(4-chlorophenoxy)- (CA INDEX NAME)

Relative stereochemistry.

PAGE 2-A

OMe

OS.CITING REF COUNT: 15 THERE ARE 15 CAPLUS RECORDS THAT CITE THIS

RECORD (15 CITINGS)

REFERENCE COUNT: 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 37 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1999:672842 CAPLUS Full-text

DOCUMENT NUMBER: 131:317743

TITLE: Drug screening with non-endogenous, constitutively

activated human serotonin receptors and small molecule

modulators thereof

INVENTOR(S): Behan, Dominic P.; Chalmers, Derek T.; Foster, Richard

J.; Glen, Robert C.; Lawless, Michael S.; Liaw, Chen W.; Liu, Qian; Russo, Joseph F.; Smith, Julian R.;

Thomsen, William J.

PATENT ASSIGNEE(S): Arena Pharmaceuticals, Inc., USA; Tripos, Inc.

SOURCE: PCT Int. Appl., 142 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 17

PATENT INFORMATION:

PATENT NO.	KIND DATE				APPLICATION NO.						DATE					
WO 9952927													19990414 <			
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	EE,															
	, KG,															
	, MX,															
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RW: AT										CD	TE	тт	ттт	мс	NIT	
	SE															
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AU 9937466			Α	1	19991	101	I	AU 19	99-3	7466			19	9904	14 <	
AU 764766			В2		2003	0828										
US 6107324			Α	2	20000	822	Ţ	JS 19	99-2	9207	1		19	9904	14 <	
US 6140509			Α	2	20001	1031	Ţ	JS 19	99-2	9206	9		19990414 <			
EP 1071701													19990414 <			
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JP 2003514	763		T	2	20030)422		TP 20	00-5	4348	3		19	9904	14 <	
MX 2000010	060		Α	2	20040)423	N	1X 20	00-1	0060			20	0010	13 <	
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AU 2004203	102		A1		2004	0729		AU 2	004-	2031	02			0040		
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AU 2007202	139		A1		2007	0531		AU 2	007-	2021	39		2	0070	510	
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ASSIGNMENT HIST	ORY F	OR U	S PA	TENT	AVA	ILAB										
					101	2100		-~								

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 131:317743

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);

AB Disclosed herein are non-endogenous, constitutively activated forms of the human 5-HT2A and human 5-HT2C receptors and uses of such receptors to screen candidate compds. Further disclosed herein are candidate compds. identified by the screening method which act at the 5HT2A receptors. Yet further disclosed is a new class of compds. which act at the 5HT2A receptors. IT 247038-30-89

BIOL (Biological study); PREP (Preparation); USES (Uses)
(drug screening with non-endogenous, constitutively activated human serotonin receptors and small mol. modulators thereof)
RN 247038-30-8 CAPLUS

CN Urea, N-[3-(4-bromo-1-methyl-1H-pyrazol-3-yl)phenyl]-N'-[(4-fluorophenyl)methyl]- (CA INDEX NAME)

$$\stackrel{\text{Me}}{\longrightarrow} \stackrel{\text{N}}{\longrightarrow} \stackrel{\text{N}}{$$

OS.CITING REF COUNT: 19 THERE ARE 19 CAPLUS RECORDS THAT CITE THIS

RECORD (20 CITINGS)

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 38 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1999:659365 CAPLUS $\underline{\text{Full-text}}$

DOCUMENT NUMBER: 131:271873

TITLE: Preparation of pyrazoles and triazoles as inhibitors

of cytokine production

INVENTOR(S): Ba Maung, Nwe Y.; Basha, Anwer; Djuric, Stevan W.;

Gubbins, Earl J.; Luly, Jay R.; Tu, Noah P.; Madar, David J.; Warrior, Usha; Wiedeman, Paul E.; Zhou, Xun;

Wagenaar, Frank L.; Sciotti, Richard J.

PATENT ASSIGNEE(S): Abbott Laboratories, USA SOURCE: PCT Int. Appl., 319 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	NO.			KIN	D i	DATE		APPLICATION NO.						DATE			
WO 9951		A1	A1 19991014			V	WO 1999-US7766					19990408 <					
W:	ΑE,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	ВG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	
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	JP,	ΚE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	
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	CI,	CM,	GA,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG						
CA 2327	185			A1	1	L9991	1014		CA 19	99-2	3271	85		19	9904	> 80	
AU 9933	879			Α	1	L9991	1025	I	AU 19	99-3	3879			19	9904	> 80	
EP 1068	187			A1	2	20010)117	E	EP 19	99-9	1534	1		19	9904	> 80	
R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	PT,	IE, FI	
JP 2002	JP 2002510679			T	20020409			Ċ	JP 2000-542301					19990408 <			
MX 2000009837 A				2	20010	0405	MX 2000-9837				20001006 <						

PRIORITY APPLN. INFO.:

US 1998-56996 WO 1999-US7766 A 19980408 <--W 19990408

<--

OTHER SOURCE(S): MARPAT 131:271873

GΙ

$$\begin{array}{c|c}
R^2 & R^3 \\
R^4 & R^5 \\
R^5 & R^5
\end{array}$$

$$^{\text{F3C}}$$
 $^{\text{N}}$ $^{\text{NH-CO}}$ $^{\text{II}}$

Title compds. [I; R1 = H, NH2, OCONH2, CN, NO2, OH, CO2H, F, C1, Br, I, aryl, perfluoroalkyl, hetercyclyloxy, hetercyclylsulfonyl; R2 = H, alkyl cycloalkyl, alkylcarbonyl, heterocyclyl; R3 = H, NH2, OCONH2, CN, NO2, OH, CO2H, F, C1, Br, I, aryl, perfluoroalkyl, hetercyclyloxy, hetercyclylsulfonyl; R4 and R5 are independently selected from H, alkyl, alkoxy, halo, perfluoroalkyl, CN, heterocycle; E = LB; B = alkyl, alkenyl, alkynyl; L = N:N, N:CH, CH:N, ON:CH, O, CO, NH, NHCO, NHSO2, NHCH2, alkenylene; Q = benzene ring with 2, 3, or 4 substituted E, heterocycle; Z = C; R2Z = N], E, Z isomers, stereoisomers, pharmaceutical acceptable salts, and prodrugs are prepared and tested as cytokine production inhibitors and are useful for treating diseases that are prevented by or ameliorated with Interleukin-2, Interleukin-4, or Interleukin-5 production inhibitors. Thus, the title compound II was prepared

IT 245746-03-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation of pyrazoles and triazoles as inhibitors of cytokine production)

RN 245746-03-6 CAPLUS

CN Propanamide, N-[4-[3,5-bis(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]-2-(4-chlorophenoxy)-2-methyl- (CA INDEX NAME)

$$F_3C$$

$$NH$$

$$CF_3$$

$$NH$$

$$CF_3$$

OS.CITING REF COUNT: 14 THERE ARE 14 CAPLUS RECORDS THAT CITE THIS

RECORD (19 CITINGS)

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 39 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1999:659233 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 131:286505

TITLE: Preparation of isoxazoloquinolinones as multidrug

resistance protein (MRP1) inhibitors

INVENTOR(S): Gruber, Joseph Michael; Kroin, Julian Stanley; Norman,

Bryan Hurst

PATENT ASSIGNEE(S): Eli Lilly and Company, USA SOURCE: PCT Int. Appl., 126 pp.

CODEN: PIXXD2

DOCUMENT TYPE: %atent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	KIN	KIND DATE				APPLICATION NO.					DATE						
WO 99512	228			A1	A1 19991014			WO 1999-US7613						19990407 <			
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	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG						
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EP 10679	928			A1	20010117 EP 1999-916456												
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HU 2001	00150	8 (A2	2	20011	1028	HU 2001-1508					19990407 <				
HU 2001						2002	0429										
JP 2002	51062	25		Τ	2	20020	0409	Ü	JP 20	00 - 5	4199	9		19	9904	07 <	
US 63690	US 6369070					20020	0409	J	JS 20	00-6	4606	2		20	0009	13 <	
MX 2000009655				A	2	20010	0316	MX 2000-9655					20001002 <				
HR 2000000646				A2	20010630			HR 2000-646					20001003 <				
NO 2000005023				Α	20001205			NO 2000-5023					20001005 <				
ORITY APPLN. INFO.:								U	IS 19	98-8	1080	P	Р	19	9804	08 <	

WO 1999-US7613 W 19990407

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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 131:286505

GΙ

AB The title compds. (I) [where R = (un) substituted amino(alkyl) or aminoethoxy, or (CH2)m'R3; m and m' = independently 0, 1, or 2; R3 = H, OH, alkoxy, amino ester, amino acid, or (un) substituted amino; R' = H, OH, or (un) substituted alkoxy] were prepared as inhibitors of 190 kDa multidrug resistance protein (MRP1) for inhibiting resistant neoplasms (14 specific neoplasm types claimed). Selected invention compds. were prepared via solution and solid phase combinatorial synthetic methods. For example, 3-(2-chloro-6-fluorophenyl)-5-methyl-4-isoxazoyl chloride was coupled with N-(5-methylisoxaz-3-oyl)-3-aminobenzylamine to form the amide followed by treatment with NaOH to yield the cyclized title compound (II). Several general procedures using substituted polystyrene resins for combinatorial preparation of title compds. were given. Representative compds. demonstrated significant reversal of MRP1 multiple drug resistance, and many compds. gave significant enhancement of oncolytic agent activities (no data). A large majority of the compds. tested were also said to have displayed a significant degree of selective inhibition of the HL60/ADR cell line over the HL60/VCR cell line in an assay for reversal of MRP1-mediated doxorubicin and vincristine resistance (no data).

IT 1101885-24-8 1101885-26-0

RL: PRPH (Prophetic)

(Preparation of isoxazoloquinolinones as multidrug resistance protein (MRP1) inhibitors)

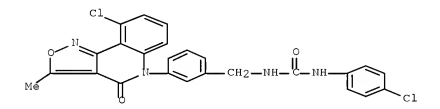
RN 1101885-24-8 CAPLUS

CN Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-[[3-(9-chloro-3-methyl-4-oxoisoxazolo[4,3-c]quinolin-5(4H)-yl)phenyl]methyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 1101885-26-0 CAPLUS

CN Urea, N-[[3-(9-chloro-3-methyl-4-oxoisoxazolo[4,3-c]quinolin-5(4H)-yl)phenyl]methyl]-N'-(4-chlorophenyl)- (CA INDEX NAME)



OS.CITING REF COUNT: 10 THERE ARE 10 CAPLUS RECORDS THAT CITE THIS

RECORD (10 CITINGS)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 40 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1999:388522 CAPLUS Full-text

DOCUMENT NUMBER: 131:80699

TITLE: Photographic couplers having UV-absorbing function and

silver halide photographic material using same

INVENTOR(S): Nakamura, Kazuaki; Chen, Zuliu; Kita, Hiroshi; Kaneko,

Yutaka

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 68 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 11160840	A	19990618	JP 1997-340814	19971127 <
PRIORITY APPLN. INFO.:			JP 1997-340814	19971127 <

OTHER SOURCE(S): MARPAT 131:80699

AB The title couplers have UV-absorbing function and molar absorption coefficient of $\geq 10,000$ at 350 nm and ≤ 100 at 420 nm in MeOH.. Cyan, magenta, and yellow couplers with specified structures and above conditions are also

claimed. A Ag halide photog. material containing ≥ 1 of the above couplers is also claimed. The photog. material shows improved processability in rapid process and provides high quality color images with excellent lightfastness.

IT 228415-90-5 228415-91-6

RL: TEM (Technical or engineered material use); USES (Uses) (photog. coupler having UV-absorbing property and specified mol. absorption coefficient for emulsion providing image with light fastness) 228415-90-5 CAPLUS

CN Dodecanamide, 2-[3-(2H-benzotriazol-2-yl)-5-(1,1-dimethylethyl)-2-hydroxyphenoxy]-N-[4-chloro-3-[[4,5-dihydro-5-oxo-1-(2,3,4,5,6-pentachlorophenyl)-1H-pyrazol-3-yl]amino]phenyl]- (CA INDEX NAME)

PAGE 1-B

RN

RN 228415-91-6 CAPLUS

CN 1-Imidazolidineacetamide, N-[5-[[2-[4-(2H-benzotriazol-2-y1)-3-hydroxyphenoxy]-1-oxotetradecyl]amino]-2-chlorophenyl]- α -(2,2-dimethyl-1-oxopropyl)-2,5-dioxo-3-(phenylmethyl)- (CA INDEX NAME)

 \sim CH2-Ph

L87 ANSWER 41 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1998:268348 CAPLUS Full-text

DOCUMENT NUMBER: 128:321662 ORIGINAL REFERENCE NO.: 128:63765a

TITLE: Compositions and methods for treating bone deficit

conditions

INVENTOR(S): Orme, Mark W.; Baindur, Nand; Robbins, Kirk G.; et al.

PATENT ASSIGNEE(S): Zymogenetics, Inc., USA; Osteoscreen, Inc.

SOURCE: PCT Int. Appl., 215 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

		KIND DATE			APPLICATION NO.													
						19980430			WO 1997-US18864									
	W:	AL,	AM,	AU,	BB,	BG,	BR,	CA,	CN,	CZ,	EE,	FΙ,	GE,	HU,	IL,	IS,	JP,	
		KG,	KP,	KR,	LK,	LR,	LT,	LV,	MD,	MG,	MK,	MN,	MX,	NO,	NZ,	PL,	RO,	
		SG,	SI,	SK,	TR,	TT,	UA,	US,	UΖ,	VN,	AZ,	BY,	KΖ,	RU,	ΤJ,	TM		
	RW:	GH,	ΚE,	LS,	MW,	SD,	SZ,	UG,	ZW,	AT,	BE,	CH,	DE,	DK,	ES,	FI,	FR,	
		GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	
		GN,	ML,	MR,	NE,	SN,	TD,	TG										
US 5990169				A	-	19991	1123	US 1997-806771					19970226 <					
US	US 6153631				A	2	20003	1128	US 1997-806768					19970226 <				
US	62519			В1	2	2001	0626	US 1997-806768 US 1997-806769					19970226 <					
US			A	-	1999(0706	US 1997-808743					19970228 <						
					A		19990	0713							19	9702	28 <	
US	US 5948776				A	-	1999(0907	J	JS 19	97-8	0873	9		19	9702	28 <	
						-	1999:	1130	Ţ	JS 19	97-8	0874	4		19	9702	28 <	
US	63425	514			В1	2	20020	0129	US 1997-808741						19	9702	28 <	
	59655	573			A	19991012								19970306 <				
AU	97498	389			A	-	19980	0515	AU 1997-49889					19971023 <				
									EP 1997-912787									
																	PT,	
		IE,	FΙ															
JP	20015	51045	50		Τ	2	2001	0731	·	JP 19	98-5	1952	9		19	9710	23 <	
US	66496	531			В1	2	2003	1118	JP 1998-519529 US 1999-297188						19	9911	19 <	
RIORITY	Y APPI	I. N.	NFO.	:					J	JS 19	96-7	3587	0	A	.2 19	9610	23 <	
									1	US 19	96-7	73581	73	P	12 19	9610	23 <	
									1	US 19	96-7	7358	74	P	12 19	9610	23 <	
									1	US 19	96-7	7358	76	P	12 19	9610	23 <	
									1	US 19	96-7	73588	31	I	12 19	9610	23 <	

Ţ	JS 1996-736220	A2	19961023 <
Ţ	JS 1996-736221	A2	19961023 <
Ţ	JS 1996-736222	A2	19961023 <
Ţ	JS 1996-736228	A2	19961023 <
Ţ	JS 1996-736318	A2	19961023 <
Ţ	JS 1996-736319	A2	19961023 <
Ī	√O 1997-US18864	W	19971023

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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 128:321662
GI

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 II

AΒ Compds. containing 2 covalently linked aromatic systems, i.e. Ar1LAr2 [I; Ar1, Ar2 = (un) substituted Ph, naphthyl, or 5- or 6-membered aromatic heterocyclyl; L = linker (atoms or covalent bond per se) so as to space the aromatic systems at a distance of 1.5-15 Å] are effective in treating conditions associated with bone deficits. The compds. can be administered to vertebrate subjects alone or in combination with addnl. agents that promote bone growth or that inhibit bone resorption. They can be screened for activity prior to administration by assessing their ability to effect the transcription of a reporter gene coupled to a promoter associated with a bone morphogenetic protein and/or their ability to stimulate calvarial growth in model animal systems. A variety of compds. were prepared and/or tested by high-throughput screening. For instance, title compound II was prepared by condensation of 2-chloro-5-(trifluoromethyl)pyridine with ethylenediamine in the presence of EtN(Pr-iso)2 at reflux. At 5-50 μg/kg/day in ovariectomized rats, II stimulated bone growth with volume increases of 21-71% observed In a calvarial bone growth assay, another compound I induced a 4-fold increase in width of new calvarial bone vs. controls.

IT 206983-31-5

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(preparation and/or use of linked aromatic and heteroarom. compds. for treating $% \left(1\right) =\left(1\right) \left(1\right) +\left(1\right) \left(1\right) \left(1\right) +\left(1\right) \left(1\right)$

bone deficit conditions)

RN 206983-31-5 CAPLUS

CN Acetamide, 2-[2,3-dichloro-4-(2-methylene-1-oxobutyl)phenoxy]-N-[4-(6-methyl-2-benzothiazolyl)phenyl]- (CA INDEX NAME)

$$\begin{array}{c} \text{O} \quad \text{CH}_2 \\ \text{NH} \quad \text{C} \quad \text{CH}_2 \quad \text{O} \quad \text{C}_1 \\ \end{array}$$

OS.CITING REF COUNT: 28 THERE ARE 28 CAPLUS RECORDS THAT CITE THIS

RECORD (42 CITINGS)

REFERENCE COUNT: 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 42 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1998:250738 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 128:294606

ORIGINAL REFERENCE NO.: 128:58387a,58390a

TITLE: Preparation of aniline derivatives having

antihyperglycemic activity

INVENTOR(S): Bierer, Donald E.; Dubenko, Larisa G. PATENT ASSIGNEE(S): Shaman Pharmaceuticals, Inc., USA

SOURCE: U.S., 41 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5741926	A	19980421	US 1997-799745	19970212 <
PRIORITY APPLN. INFO.:			US 1997-799745	19970212 <
ACCIONIMENTE LITOTODIA DOD	770 DAMES	10 3 T T 3 T T 3 T T T T	THE FOLIO DECDERS BODIES	

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 128:294606

GI

AB The title compds. [I; R1-R5 = H, halo, C1-6 alkyl, etc.; R7-R10 = H, halo, Ph, etc.; A = C(O), CH2; B = NH, O, S], useful for the treatment of insulin-dependent diabetes mellitus (IDDM or Type I) and non-insulin dependent diabetes mellitus (NIDDM or Type II), were prepared Thus, treatment of anthranilic acid with bromoacetyl bromide in DMF and dioxane followed by reaction of the resulting 2-[(2-bromoacetyl)amino]benzoic acid with o-fluoroaniline in DMF afforded the title compound II which showed stimulatory effect (128% basal) on 2-deoxy-D-glucose uptake in 3T3-L1 adipocytes in the absence of insulin.

Т

IT 195393-81-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of aniline derivs. having antihyperglycemic activity)

RN 195393-81-8 CAPLUS

CN Acetamide, 2-[(2-fluorophenyl)amino]-N-[4-(2H-tetrazol-5-yl)phenyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD

(4 CITINGS)

REFERENCE COUNT: 53 THERE ARE 53 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 43 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1997:574584 CAPLUS Full-text

DOCUMENT NUMBER: 127:212475

ORIGINAL REFERENCE NO.: 127:41189a,41192a

TITLE: N-(Heterocyclylaryl)hydrazine derivative for a

principal color developer, silver halide photographic

light-sensitive material and imaging method

INVENTOR(S):

Okawa, Atsuhiro; Makuta, Toshiyuki; Taguchi, Toshiki

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 82 pp.

CODEN: JKXXAF

DOCUMENT TYPE: %atent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 09211818 US 5851749	 А А	19970815 19981222	JP 1996-331409 US 1996-757730	19961128 < 19961126 <
PRIORITY APPLN. INFO.:			JP 1995-334183 A	. 19951130 <
ASSIGNMENT HISTORY FOR OTHER SOURCE(S): GI		T AVAILABLE 127:212475	IN LSUS DISPLAY FORMA	T

AΒ The title compds. [I; Z1 = acyl, CONH2, alkoxycarbonyl, aryloxycarbonyl, R1SO2, C(X):NR2; wherein R1 = alkyl, alkenyl, alkynyl, aryl, heterocyclyl; X = OR3, NR4R5; R2, R4, R5 = H, alkyl, alkenyl, alkynyl, aryl, heterocyclyl; R3 = same as R1; or R2 and R3, or R4 and R5 are bonded together to form a ring; Q1 = a group of nonmetal atoms necessary to form a 5- or 6-membered ring together with the C atom; Q2 = heterocyclyl; Y = substitutable group; m = 1,2; n = 0-3] (e.g. II) are prepared An imaging method involves development of an imagewise-exposed silver halide photog. light-sensitive material in the presence of above color developer I, in particular with a processing liquid containing above color developer I. A silver halide photog. light-sensitive material comprises at least one hydrophilic colloidal layer containing above color developer I formed on a support. Another imaging method involves development of the latter photog. material (1) by heat-treatment at $50-200^{\circ}$ or (2) in a solution These compds. provide new principal developers which form dyes excellent in coloration during development and give images of good coloration and stability and stable in hue even when couplers substituted at the coupling position are used. 194790-64-2 ΙT

RL: TEM (Technical or engineered material use); USES (Uses) (photog. color developer; N-(heterocyclylaryl)hydrazine derivs. for principal color developers, silver halide photog. light-sensitive material, and imaging method)

RN 194790-64-2 CAPLUS

CN Hydrazinecarboxamide, 2-[2,4-bis(1-methyl-1H-benzimidazol-2-yl)phenyl]-N-[2-chloro-5-[(dioctylamino)sulfonyl]phenyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

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L87 ANSWER 44 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1997:563089 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 127:247927

ORIGINAL REFERENCE NO.: 127:48437a,48440a

TITLE: Aniline derivatives having antihyperglycemic activity

INVENTOR(S): Bierer, Donald E.; Dubenko, Larisa G. PATENT ASSIGNEE(S): Shaman Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 129 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 9730019 19970821 WO 1997-US2289 Α1 19970213 <--W: AL, AM, AU, AZ, BA, BB, BG, BR, BY, CA, CN, CU, CZ, EE, GE, HU, IL, IS, JP, KG, KP, KR, KZ, LC, LK, LR, LT, LV, MD, MG, MK, MN, MX, NO, NZ, PL, RO, RU, SG, SI, SK, TJ, TM, TR, TT, UA, UZ, VN, YURW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG AU 9721241 19970902 AU 1997-21241 19970213 <--Α PRIORITY APPLN. INFO.: US 1996-600725 A 19960213 <--WO 1997-US2289 19970213

OTHER SOURCE(S):

MARPAT 127:247927

Ι

GI

Aniline derivs. useful as antihyperglycemic agents, pharmaceutical compns. comprising the aniline derivs., and methods for their use are described. For instance, the novel compds. I [R1-R5 = H, halo, OR11, CX3, alkyl, (CH2)nCH2OH, (CH2)nCO2R12, (CH2)nT; one and only one of R1-R5 = one of the latter 2 groups; R11, R12 = H, alkyl; X = halo; n = 0, 1; R7-R10 = H, halo, OR13, SR14, CY3, alkyl, Ph; R13, R14 = H, alkyl, Ph; Y = halo; A = CO, CH2; B = NH, O, S; T = 5-tetrazolyl] are described. The aniline derivs. are useful for the treatment of insulin-dependent and non-insulin-dependent diabetes mellitus. For instance, amidation of anthranilic acid with BrCH2COBr in DMF/dioxane (87.8% yield) and condensation of the intermediate bromo compound with o-fluoroaniline in DMF (85% yield) gave title compound II, a preferred compound At 100 mg/kg orally in diabetic db/db mice, II reduced blood glucose by 61.3 mg/dL at 27 h, vs. 116.4 mg/dL for metformin at the same dosage.

IT 195393-81-8P, 4-(Tetrazol-5-yl)-1-[[2-[(2-fluorophenyl)amino]acetyl]amino]benzene

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of aniline derivs. with antihyperglycemic activity)

RN 195393-81-8 CAPLUS

CN Acetamide, 2-[(2-fluorophenyl)amino]-N-[4-(2H-tetrazol-5-yl)phenyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS)

L87 ANSWER 45 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1996:116243 CAPLUS Full-text

DOCUMENT NUMBER: 124:260935

ORIGINAL REFERENCE NO.: 124:48343a,48346a

TITLE: Synthesis and antimicrobial activities of some new

benzimidazoles, Part I

AUTHOR(S): El-Sherief, H. A.; El-Ezbawy, S. R.; Mahmoud, A. M.;

Sarhan, Abd El-Wareth A. O.

CORPORATE SOURCE: Faculty Science, Assiut University, Assiut, Egypt
SOURCE: Bulletin of the Faculty of Science, Assiut University,

B: Chemistry (1995), 24(1), 111-23

CODEN: BFSAE6; ISSN: 1010-2671

PUBLISHER: Assiut University

DOCUMENT TYPE: Journal LANGUAGE: English

AB Reaction of Et p-(2-benzimidazolyl)phenoxyacetate (1) with aromatic amines gave the corresponding acetanilides. Reaction of 1 with hydrazine hydrate gave the hydrazide, which reacted with aromatic aldehydes, acetylacetone, Et acetoacetate, CS2, etc. Antibacterial activity of several derivs. was determined

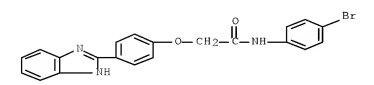
IT 175028-44-1P

RL: SPN (Synthetic preparation); PREP (Preparation)

(synthesis and antimicrobial activities of benzimidazole derivs.)

RN 175028-44-1 CAPLUS

CN Acetamide, 2-[4-(1H-benzimidazol-2-yl)phenoxy]-N-(4-bromophenyl)- (CA INDEX NAME)



OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

L87 ANSWER 46 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1994:148835 CAPLUS Full-text

DOCUMENT NUMBER: 120:148835

ORIGINAL REFERENCE NO.: 120:25989a,25992a

TITLE: Photographic coupler and silver halide color

photographic material

INVENTOR(S): Mizukawa, Yuki; Motoki, Masuji; Sato, Tadahisa;

Takahashi, Osamu

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan

SOURCE: Eur. Pat. Appl., 126 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
	EP 571959	A2	19931201	EP 1993-108447		19930525 <
	EP 571959	A3	19940316			
	EP 571959	В1	19970409			
	R: DE, FR, GB,	NL				
	JP 06043611	A	19940218	JP 1992-234120		19920811 <
	EP 688774	A1	19951227	EP 1995-112304		19930525 <
	EP 688774	B1	19991222			
	R: DE, FR, GB,	NL				
	US 5451501	A	19950919	US 1993-67111		19930526 <
	JP 08109181	A	19960430	JP 1995-158770		19950602 <
	US 5532377	A	19960702	US 1995-467833		19950606 <
PRIO	RITY APPLN. INFO.:			JP 1992-157405	A	19920526 <
				JP 1992-234120	A	19920811
<						
				EP 1993-108447	A3	19930525 <
				US 1993-67111	A3	19930526 <

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): CASREACT 120:148835

GΙ

AB A 1H-pyrazolo[1,5-b][1,2,4]triazole magenta coupler having a t-alkyl group at the position-6 and an amido group-substituted Ph group at the position-2 is disclosed having the formula I [R1 = tert-alkyl; R2, R3 = H, substituent; Y = H, halogen, aryloxy; A, B = CO, SO2; n = 0, 1; R4 = H, alkyl, aryl; R5

= alkyl, aryl, alkoxy, alkylamino, arylamino; R4-R5 may be combined to form a ring]. There is also disclosed a Ag halide color photog. material containing the same. The couplers provide excellent latent image stability, cause no lowering of sensitivity, and have excellent color developability. 152828-07-49

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and use of, as photog. magenta coupler)

RN 152828-07-4 CAPLUS

ΙT

CN Hexanamide, N-[4-[7-chloro-6-(1,1-dimethylethyl)-3H-pyrazolo[1,5-b][1,2,4]triazol-2-yl]phenyl]-2-[2-chloro-4-(1,1,3,3-tetramethylbutyl)phenoxy]- (CA INDEX NAME)

OS.CITING REF COUNT: 6 THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS)

L87 ANSWER 47 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1993:29854 CAPLUS Full-text

DOCUMENT NUMBER: 118:29854

ORIGINAL REFERENCE NO.: 118:5361a,5364a

TITLE: Silver halide color photographic material

INVENTOR(S): Shimada, Yasuhiro; Ishii, Yoshio PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 37 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO	•	DATE
JP 04190348	A	19920708	JP 1990-322051		19901126 <
US 5272051	A	19931221	US 1991-788432		19911106 <
PRIORITY APPLN. INFO.:			JP 1990-322051	A	19901126 <
ASSIGNMENT HISTORY FOR	US PATEN	T AVAILABLE	IN LSUS DISPLAY	FORMAT	
GI					

AB In the title material comprising a support having thereon one or more red-sensitive silver halide emulsion layers, at least one of the said red-sensitive silver halide emulsion layers contains a cyan coupler represented by general structure I. For I, R = a substituent group; Z1 = an electron-attracting substituent group which is not released upon reaction with an oxidized aromatic primary amine developing agent; Z2 = an electron-attracting substituent group; T = H or a group to be released upon reaction with an oxidized aromatic primary amine developing agent. The title material provides excellent color reproduction

IT 145130-89-8

RL: TEM (Technical or engineered material use); USES (Uses) (photog. cyan coupler)

RN 145130-89-8 CAPLUS

CN 1H-Imidazo[1,2-b]pyrazole-7-carboxamide,

3-[[2-butoxy-4-(1,1,3,3-tetramethylbutyl)phenyl]thio]-6-[4-[[2-(2-chlorophenoxy)-1-oxotetradecyl]amino]phenyl]-2-cyano- (CA INDEX NAME)

L87 ANSWER 48 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1992:521410 CAPLUS Full-text

DOCUMENT NUMBER: 117:121410

ORIGINAL REFERENCE NO.: 117:20933a,20936a

TITLE: Photographic material using diffusion-resisting dye INVENTOR(S): Ohashi, Hirobumi; Kaqawa, Nobuaki; Kaquchi, Hiroyuki

PATENT ASSIGNEE(S): Konica K. K., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 20 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	JP 04046337	A	19920217	JP 1990-154834	19900613 <
	JP 2852455	B2	19990203		
PRIO	RITY APPLN. INFO.:			JP 1990-154834	19900613 <
GT					

$$\begin{array}{c}
\begin{array}{c}
\text{CN} \\
\text{N} \\
\text{N}
\end{array}$$

$$\begin{array}{c}
\text{L}^{1} + \text{L}^{2} - \text{L}^{3} \\
\text{MO}
\end{array}$$

$$\begin{array}{c}
\text{CN} \\
\text{N}
\end{array}$$

AB The title photog. material contains a dispersion of ≥ 1 oil-soluble dye I [R1, R2 = alkyl, aryl, aralkyl, alkenyl, heterocyclyl; L1-3 = methine; n = 0-2; M = H, monovalent metal atom] in its photog. component layers. The photog. material shows reduced fog and improved long-term stability.

IT 142966-19-6

RL: TEM (Technical or engineered material use); USES (Uses) (photog. materials containing)

RN 142966-19-6 CAPLUS

CN Octanamide,

2-[2-chloro-4-(1,1-dimethylpropyl)phenoxy]-N-[4-[4-[[1-[4-[[2-

[2-chloro-4-(1,1-dimethylpropyl)phenoxy]-1-oxooctyl]amino]phenyl]-3-cyano-1,5-dihydro-5-oxo-4H-pyrazol-4-ylidene]methyl]-3-cyano-5-hydroxy-1H-pyrazol-1-yl]phenyl]- (CA INDEX NAME)

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L87 ANSWER 49 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1991:460720 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 115:60720

ORIGINAL REFERENCE NO.: 115:10299a,10302a

TITLE: High-contrast high-sensitivity rapidly processable

photographic material

INVENTOR(S): Hirano, Shigeo; Ichijima, Yasushi; Deguchi, Hisayasu

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 64 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 02213 8 38	A	19900824	JP 1989-35871	19890215 <
PRIORITY APPLN. INFO.:			JP 1989-35871	19890215 <

AB The title photog. material utilizes ≥ 1 A-(L1)v-B-(L2)w-FA [A = group releasing (L1)v-B-(L2)w-FA on reacting with oxidized developing agent; L1 = group releasing B-(L2)w-FA following its separation from A; B = group releasing (L2)w-FA on reacting with oxidized developing agent; L2 = group releasing FA following its separation from B; FA = nucleating agent, development accelerator; v,w = 0, 1]. The above compound releases a fogging agent on development accelerator in an imagewise manner and has a high maximum d.

IT 134889-42-2

RL: USES (Uses)

(photog. additive, high maximum d. materials using)

RN 134889-42-2 CAPLUS

CN Benzoic acid, 4-chloro-3-[[2-[[6-[(dodecylamino)carbony1]-7-[[[[4-(2-formylhydrazino)pheny1]amino]carbony1]-1H-benzotriazol-1-y1]-1,2,3,4-tetrahydro-1,4-methanonaphthalen-5-y1]oxy]-3-(4-methoxypheny1)-1,3-dioxopropy1]amino]-, 2-(dodecyloxy)-1-methyl-2-oxoethyl ester (9CI) (CA INDEX NAME)

L87 ANSWER 50 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1991:81825 CAPLUS Full-text

DOCUMENT NUMBER: 114:81825

ORIGINAL REFERENCE NO.: 114:13973a, 13976a

TITLE: Substituted N-(heterocyclic-substituted phenyl)-N'-benzylureas as pesticides

INVENTOR(S): Carney, Robert L.; Gruber, John M.; Lui, Alfred S. T.

PATENT ASSIGNEE(S): Sandoz A.-G., Switz.

SOURCE: U.S., 16 pp. Cont.-in-part of U.S. Ser. No. 187,164.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 4950678	A	19900821	US 1989-386333	19890727 <
PRIORITY APPLN. INFO.:			US 1986-840814	B2 19860318 <
			US 1987-12577	B2 19870209 <
			US 1988-187164	A2 19880428 <

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 114:81825

GΙ

$$\begin{array}{c} X1 \\ X2 \\ X3 \\ X2 \end{array}$$
 CONHC (Y) NH $\begin{array}{c} X6 \\ X5 \\ X4 \\ \end{array}$ R2

AB Title compds. I (X1,X2,X3,X5 = H, halo, C1-8 alkyl, C1-8 alkoxy; X4 = H, halo, (substituted) (halo) C1-8 alkyl, etc.; R6 = H, halo, C1-8 alkyl, etc.; Y = O, S; A = R4C, R4 = H, halo, (halo) C1-8 alkyl, (substituted) (halo) C1-8 alkoxy, (halo) C1-8 alkylthio, etc.; B = R3C, R3 = R4; R1 = R4; R2 = R3) and salts thereof, useful as pesticides in particular acaricides and insecticides (no data), are prepared 2,6-F2C3H3CONCO was added to 4-(4-chloro-1-pyrazolyl)aniline (preparation given) in CH2C12 to give I (X1 = X2 = F, X3 - X6 = R1 = H, R2 = C1, Y = O, A = N, B = CH).

IT	112736-96-6P	112736-97-79	112736-98-8P
	112736-99-99	112737-00-52	112737-01-6P
	112737-02-79	112737-03-89	112737-04-9P
	112737-05-0P	112737-06-19	112737-07-20
	112737-08-39	112737-09-42	112737-10-79
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	112737-14-1P	112737-15-2P	112737-16-39
	112737-17-4P	112737-18-5P	112737-19-69
	112737-20-92	112737-21-0P	112737-22-1P
	112737-23-29	112737-24-3P	112737-25-4P
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	112737-39-0P	112737-40-39	112737-41-4P
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	112737-45-8P	112737-46-9P	112737-47-0P
	112737-48-1P	112737-50-5P	112737-51-6P
	112737-52-7P	112737-53-88	112737-54-99

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             112737-61-8P
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                              112737-87-8P
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                              112738-02-0P
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                              131778-69-39
               131778-71-79
                              131778-72-89
131778-70-62
131778-73-92
               131778-74-09
                              131778-75-1P
               131778-77-3P
131778-76-22
                              131778-78-4P
131778-79-52
               131778-80-8P
                              131778-81-9P
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                              131778-98-8P
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131797-34-7P
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RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as insecticide)

RN 112736-96-6 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112736-97-7 CAPLUS

CN Benzamide, N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-3,5-dimethylphenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\underbrace{ \begin{array}{c} F \\ C-NH-C-NH \\ \end{array} }_{F} \underbrace{ \begin{array}{c} Me \\ N\\ \end{array} }_{C1} \underbrace{ \begin{array}{c} C1 \\ \end{array} }_{C1} \underbrace{ \begin{array}{c$$

RN 112736-98-8 CAPLUS

CN Benzamide, N-[[[4-(4-bromo-1H-pyrazol-1-y1)-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

RN 112736-99-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3,4-dibromo-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-00-5 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F3C$$

$$C1$$

$$NH-C-NH-C$$

$$C1$$

$$C1$$

$$C1$$

$$C1$$

RN 112737-01-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-(4-chloropheny1)-1H-pyrazol-1-

yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-02-7 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-03-8 CAPLUS

CN Benzamide, N-[[[4-[3-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-04-9 CAPLUS

CN Benzamide, N-[[[4-(4-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & &$$

RN 112737-05-0 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-(1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} C1 & & & \\ N & & & \\ N & & & \\ \end{array}$$

RN 112737-06-1 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(4-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-07-2 CAPLUS

CN Benzamide, N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-08-3 CAPLUS

CN Benzamide, N-[[[4-[3,4-bis(trifluoromethyl)-1H-pyrazol-1-y1]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F_{3}C$$

$$NH - C - NH - C$$

$$F_{3}C$$

RN 112737-09-4 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3-chloro-1H-pyrazol-1-

yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$C1 \longrightarrow NH-C-NH-C$$

RN 112737-10-7 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3,4-dibromo-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-11-8 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-12-9 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

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RN 112737-13-0 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-14-1 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-15-2 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F3C \longrightarrow NH-C-NH-C$$

RN 112737-16-3 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(1,1-dimethylethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-17-4 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3,4-dichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$C1 \longrightarrow NH - C - NH - C \longrightarrow F$$

RN 112737-18-5 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-[4-chloro-3-(trifluoromethyl)-1H-pyrazol-1-

yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F3C \longrightarrow NH-C-NH-C$$

RN 112737-19-6 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F_{3}C$$

$$C_{1}$$

$$C_{2}$$

$$C_{3}$$

$$C_{4}$$

$$C_{5}$$

$$C_{1}$$

$$C_{1}$$

$$C_{1}$$

$$C_{1}$$

$$C_{2}$$

$$C_{3}$$

$$C_{4}$$

$$C_{5}$$

$$C_{5}$$

$$C_{7}$$

$$C_{1}$$

$$C_{7}$$

$$C_{$$

RN 112737-20-9 CAPLUS

CN Benzamide, N-[[[4-[3,5-bis(trifluoromethyl)-1H-pyrazol-1-y1]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F_3C \xrightarrow{\qquad \qquad N \\ CF_3} C1 \xrightarrow{\qquad \qquad NH-C-NH-C} F$$

RN 112737-21-0 CAPLUS

CN Benzamide, N-[[[4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-22-1 CAPLUS

CN Benzamide, N-[[[3-chloro-4-(4-chloro-1H-pyrazol-1-

yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 112737-23-2 CAPLUS

CN Benzamide, N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-y1]-3-(trifluoromethyl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-24-3 CAPLUS

CN Benzamide, N-[[[4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]-3-(trifluoromethyl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F3C \xrightarrow{N} NH C NH C F3$$

RN 112737-25-4 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-26-5 CAPLUS

CN Benzamide, N-[[[4-[4-bromo-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-27-6 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3,4,5-tribromo-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-28-7 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-[3-chloro-5-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$C1 \longrightarrow NH - C - NH - C \longrightarrow F$$

$$CF_3 \qquad C1$$

RN 112737-29-8 CAPLUS

CN Benzamide, N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-2-fluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-30-1 CAPLUS

CN Benzamide,

 $\begin{tabular}{ll} $N-[[[5-chloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-2-fluorophenyl]amino]carbonyl]-2,6-difluoro-(CA INDEX NAME) \\ \end{tabular}$

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 112737-31-2 CAPLUS

CN Benzamide, N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-3-methylphenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

RN 112737-32-3 CAPLUS

CN Benzamide,

N-[[[5-chloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-2-methylphenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-33-4 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4-(2,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

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RN 112737-34-5 CAPLUS

CN Benzamide, N-[[[4-[4-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

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RN 112737-35-6 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4-chloro-3-(4-chlorophenyl)-5-methoxy-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-37-8 CAPLUS

CN Benzoic acid, 2-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-5-[[[(2,6-difluorobenzoyl)amino]carbonyl]amino]-, methyl ester (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ &$$

RN 112737-38-9 CAPLUS

CN Benzamide, N-[[[3-chloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-39-0 CAPLUS

CN Benzamide,

N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3-chlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & &$$

RN 112737-40-3 CAPLUS

CN Benzamide, N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3- (trifluoromethyl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 112737-41-4 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(2,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-42-5 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(3,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-43-6 CAPLUS

CN Benzamide, N-[[[4-[3-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & &$$

RN 112737-44-7 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(4-fluorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-45-8 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-[3-[3-(trifluoromethyl)phenyl]-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-46-9 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(5-chloro-2-thienyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-47-0 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(2-naphthalenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 112737-48-1 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(4-cyano-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-50-5 CAPLUS

CN Benzamide, N-[[[3-chloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-51-6 CAPLUS

CN Benzamide,

N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3-chlorophenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 112737-52-7 CAPLUS

CN Benzamide, N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3-(trifluoromethyl)phenyl]amino]carbonyl]-2-fluoro-(CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 112737-53-8 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(2,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-54-9 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(3,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-55-0 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(4-fluorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-56-1 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-[3-[3-(trifluoromethyl)phenyl]-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-59-4 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-

1H-pyrazol-1-yl]phenyl]amino]carbonyl]-5-fluoro- (CA INDEX NAME)

$$F3C$$

$$C1$$

$$C1$$

$$NH-C-NH-C$$

$$C1$$

$$C1$$

RN 112737-60-7 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-6-fluoro- (CA INDEX NAME)

RN 112737-61-8 CAPLUS

CN Benzamide, 2,6-dichloro-N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F_3C$$

$$C_1$$

RN 112737-62-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-63-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(4-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

RN 112737-64-1 CAPLUS

CN Benzamide, N-[[[4-[3,4-bis(trifluoromethyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

$$F_{3}C$$

$$NH-C-NH-C$$

$$F_{3}C$$

RN 112737-65-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-66-3 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[4-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

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RN 112737-67-4 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[4-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & & & & & & & & & & & & & \\ & & & & & & & & & & & & & & & \\ & & & & & & & & & & & & & \\ & & & & & & & & & & & & \\ & & & & & & & & & & & & \\ & & & & & & & & & & & \\ & & & & & & & & & & \\ & & & & & & & & & & \\ & & & & & & & & & \\ & & & & & & & & & \\ & & & & & & & & & \\ & & & & & & & & \\ & & & & & & & & \\ & & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & \\ & & \\ & & \\ & \\ & & \\ & \\ & & \\ & \\ & \\ & & \\ \\ & \\ & \\ \\ & \\ & \\ \\ & \\ \\ & \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\$$

RN 112737-68-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F3C \longrightarrow NH-C-NH-C$$

RN 112737-69-6 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[3-(1,1-dimethylethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-70-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-71-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[4-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-72-1 CAPLUS

CN Benzamide, N-[[[4-[3,5-bis(trifluoromethyl)-1H-pyrazol-1-y1]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

$$F3C \longrightarrow NH-C-NH-C$$

RN 112737-73-2 CAPLUS

CN Benzamide, N-[[[4-[4-bromo-3-(4-chlorophenyl)-lH-pyrazol-l-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

RN 112737-74-3 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3,4,5-tribromo-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-75-4 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-chloro-5-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$C1 \longrightarrow NH-C-NH-C$$

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} C1 & & C1 \\ \hline \\ C1 & & NH-C-NH-C \\ \hline \\ C2 & & NH-C-NH-C \\ \hline \\ C3 & & NH-C-NH-C \\ \hline \\ C4 & & NH-C-NH-C \\ \hline \\ C5 & & NH-C-NH-C \\ \hline$$

RN 112737-77-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-3,5-dimethylphenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-78-7 CAPLUS

CN Benzamide,

N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3-chlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

RN 112737-79-8 CAPLUS

CN Benzamide, N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3-(trifluoromethyl)phenyl]amino]carbonyl]-2-chloro-(CA INDEX NAME)

RN 112737-80-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-(2-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-81-2 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[3-(2,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-82-3 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[3-(3,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-83-4 CAPLUS

CN Benzamide, N-[[[4-[3-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

RN 112737-84-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-5-methylphenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 112737-85-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-(4-fluorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 112737-87-8 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[3-(5-chloro-2-thienyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\bigcup_{C_1}^{\circ}\bigcup_{NH}^{\circ}\bigcup_{$$

RN 112737-88-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-(2-naphthalenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

RN 112737-89-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F3C \longrightarrow NH-C-NH-C$$

RN 112737-90-3 CAPLUS

CN Benzamide, N-[[[4-[5-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

RN 112737-91-4 CAPLUS

CN Benzamide, N-[[[4-[5-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-92-5 CAPLUS

CN Benzamide, N-[[[4-[5-(4-bromophenyl)-4-chloro-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

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 \int_{1}

CN Benzamide, N-[[[3,5-dichloro-4-[3-(4-methylphenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-94-7 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-(4-methylphenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

RN 112737-96-9 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3-chloro-1H-indazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-97-0 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(5,6-dichloro-1H-benzimidazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112738-00-8 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(5,6-dichloro-1H-benzotriazol-l-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112738-01-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(5,6-dichloro-1H-benzimidazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112738-02-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-(5,6-dichloro-1H-benzimidazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112738-03-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(5-chloro-1H-benzotriazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112738-04-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(5,6-dichloro-1H-benzotriazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112738-06-4 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-(2H-indazol-2-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112738-07-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(2H-indazol-2-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & & & & & & & & & & & & & \\ & & & & & & & & & & & & & & & \\ & & & & & & & & & & & & & & \\ & & & & & & & & & & & & & \\ & & & & & & & & & & & & \\ & & & & & & & & & & & \\ & & & & & & & & & & & \\ & & & & & & & & & & \\ & & & & & & & & & & \\ & & & & & & & & & \\ & & & & & & & & & \\ & & & & & & & & \\ & & & & & & & & \\ & & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & \\ & & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & \\ & & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ & \\ \\ &$$

RN 112762-62-6 CAPLUS

CN Benzamide, N-[[[4-[3,5-bis(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112762-63-7 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3,4-dichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112762-64-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112762-65-9 CAPLUS

CN Benzamide,

2-chloro-N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-3-(trifluoromethyl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112762-66-0 CAPLUS

CN Benzamide,

2-chloro-N-[[[4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]-3-(trifluoromethyl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112762-67-1 CAPLUS

CN Benzamide, N-[[[4-[5-(4-bromophenyl)-4-chloro-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

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RN 112785-82-7 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(2-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\bigcap_F \bigcap_{C-NH-C-NH} \bigcap_{C1} \bigcap_{N} \bigcap_{C1} \bigcap_{N} \bigcap_{C1} \bigcap_{N} \bigcap_{C1} \bigcap_{N} \bigcap_{C1} \bigcap_{C1}$$

RN 131778-69-3 CAPLUS

CN 1H-Pyrazole-4-carboxylic acid, 1-[2,6-dichloro-4-[[[(2,6-difluorobenzoyl)amino]carbonyl]amino]phenyl]-, ethyl ester (CA INDEX NAME)

RN 131778-70-6 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4-chloro-3-(4-fluorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 131778-71-7 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F3C \longrightarrow NH - C - NH - C$$

RN 131778-72-8 CAPLUS

CN Benzamide, N-[[[4-(4-bromo-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-

difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 131778-73-9 CAPLUS

CN Benzamide,

N-[[[4-(4-bromo-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2-fluoro-(CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &$$

RN 131778-74-0 CAPLUS

CN Benzamide,

 $\label{eq:normalization} $$N-[[[4-(4-bromo-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2-chloro-(CA INDEX NAME)$$

RN 131778-75-1 CAPLUS

CN Benzamide,

N-[[[4-(4-bromo-3-fluoro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 131778-76-2 CAPLUS

CN Benzamide,

N-[[[4-(4-bromo-3-fluoro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

$$\begin{array}{c|c} F & & \\ &$$

RN 131778-77-3 CAPLUS

CN Benzamide,

N-[[[4-(4-bromo-3-fluoro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

RN 131778-78-4 CAPLUS

CN Benzamide, N-[[[4-[4-bromo-3-(4-methoxyphenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & &$$

RN 131778-79-5 CAPLUS

CN Benzamide, N-[[[4-[4-bromo-3-(4-methoxyphenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

RN 131778-80-8 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3,4-diphenyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 131778-81-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3,4-diphenyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 131778-82-0 CAPLUS

CN Benzamide, N-[[[4-[3,4-bis(4-chlorophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 131778-83-1 CAPLUS

CN Benzamide, N-[[[4-[3,4-bis(4-chlorophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

RN 131778-84-2 CAPLUS

CN Benzamide, N-[[[4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 131778-85-3 CAPLUS

CN Benzamide,

2-chloro-N-[[[4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 131778-86-4 CAPLUS

CN Benzamide, N-[[[4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]-3-(trifluoromethyl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 131778-87-5 CAPLUS
CN Benzamide,
2-chloro-N-[[[4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]-3-(trifluoromethyl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 131778-90-0 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-[4-(4-chlorophenyl)-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

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RN 131778-91-1 CAPLUS

CN Benzamide,

N-[[[2,5-dichloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 131778-92-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[2,5-dichloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 131778-93-3 CAPLUS

CN Benzamide, 2-chloro-N-[[[2,3-dichloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 131778-94-4 CAPLUS

CN Benzamide,

N-[[[2,3-dichloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & &$$

RN 131778-95-5 CAPLUS

CN Benzamide, N-[[[2,3-dichloro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 131778-96-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[2,5-dichloro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 131778-97-7 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-2-fluoro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 131778-98-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-2-fluoro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$C1 \longrightarrow NH-C-NH-C$$

RN 131778-99-9 CAPLUS

CN Benzamide,

2-chloro-N-[[[3-chloro-2-fluoro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 131779-00-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[2,3,5-trichloro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 131779-01-6 CAPLUS

CN Benzamide,

2,6-difluoro-N-[[[2,3,5-trichloro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$C1 \longrightarrow NH - C - NH - C \longrightarrow F$$

RN 131797-31-4 CAPLUS

CN Benzamide, N-[[[3-chloro-4-(2,5,6-trichloro-1H-benzimidazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 131797-32-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-(trifluoromethyl)-4-[4-(trifluoromethyl)phenyl]-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

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RN 131797-33-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[2,3-dichloro-4-(3,4,5-trichloro-1H-pyrazol-1-

yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 131797-34-7 CAPLUS

CN Benzamide, N-[[[3-chloro-2-fluoro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD

(2 CITINGS)

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 51 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1991:72228 CAPLUS Full-text

DOCUMENT NUMBER: 114:72228

ORIGINAL REFERENCE NO.: 114:12161a,12164a

TITLE: Color photographic material with wide exposure

latitude

INVENTOR(S): Hirabayashi, Shigeto; Matsuzaka, Syoji; Ohya, Yukio;

Nakayama, Tomoyuki; Hoshino, Hiroyuki

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Eur. Pat. Appl., 158 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 365348 EP 365348	A2 A3	19900425 19901114	EP 1989-310819	19891020 <
R: DE, GB JP 02110551 JP 02131234 JP 02219053	A A A	19900423 19900521 19900831	JP 1988-264591 JP 1988-286538 JP 1989-40920	19881020 < 19881111 < 19890220 <

A	19900903	JP 1989-40779		19890221 <
		JP 1988-264591	A	19881020 <
		JP 1988-286538	A	19881111
		JP 1989-40920	A	19890220
		JP 1989-40779	A	19890221
	A	A 19900903	JP 1988-264591 JP 1988-286538 JP 1989-40920	JP 1988-264591 A JP 1988-286538 A JP 1989-40920 A

OTHER SOURCE(S): MARPAT 114:72228

AB Color photog. materials having a wide exposure latitude, which can provide an image with excellent graininess and sharpness, and which have high stability under variable processing conditions, consist of blue-sensitive, green-sensitive, and red-sensitive Ag halide emulsion layers on a support and ≥1 of the color-sensitive emulsion layers has a single-layer structure. The emulsion layer having a single-layer structure comprises ≥2 types of Ag halide grains differing in average grain size and contains a development-inhibitor-releasing compound The exposure latitude of the Ag halide emulsion layer with a single-layer structure is ≥3.0.

IT 125329-20-6

RL: USES (Uses)

(photog. yellow shift coupler, color materials containing, with wide exposure latitude)

RN 125329-20-6 CAPLUS

CN Benzothiazolium,

chloro-5-[(dodecyloxy)carbonyl]phenyl]amino]carbonyl]-2-(4-methoxyphenyl)2-oxoethoxy]phenyl]-3-methyl-, iodide (9CI) (CA INDEX NAME)

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L87 ANSWER 52 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1990:468229 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 113:68229

ORIGINAL REFERENCE NO.: 113:11357a,11360a

TITLE: Rapid processing of color photographic material for

stable images

INVENTOR(S): Ezaki, Atsuo; Yoshimoto, Hiroshi

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 36 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01230045	A	19890913	JP 1988-57232	19880310 <
PRIORITY APPLN. INFO.:			JP 1988-57232	19880310 <

AB In processing an exposed color photog. material by treating with a color developing agent, bleaching, and fixing, the bleaching time plus fixing time is ≤3 min 45 s and the photog. material contains a complex having a dye or its precursor bonded directly or via a timing group to the reactive site at which reaction occurs with the oxidized primary aromatic amine developer, the dye or its precursor released above having an absorption maximum shifted to a lower wavelength prior to its release.

IT 125329-20-6

RL: USES (Uses)

(yellow photog. coupler)

RN 125329-20-6 CAPLUS

CN Benzothiazolium,

[[2, 4-bis(1, 1-dimethylpropyl)phenyl]methoxy]-2-[4-[1-[[[2-bis(1, 1-dimethylpropyl)phenyl]methoxy]]

chloro-5-[(dodecyloxy)carbonyl]phenyl]amino]carbonyl]-2-(4-methoxyphenyl)-2-oxoethoxy]phenyl]-3-methyl-, iodide (9CI) (CA INDEX NAME)

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L87 ANSWER 53 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1990:449660 CAPLUS Full-text

DOCUMENT NUMBER: 113:49660

ORIGINAL REFERENCE NO.: 113:8265a,8268a

TITLE: Silver halide color photographic material containing

shift coupler

INVENTOR(S): Ueda, Eiichi; Nakagawa, Satoshi; Shimazaki, Hiroshi

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 24 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 0 121746 0	A	19890831	JP 1988-43440	19880226 <
PRIORITY APPLN. INFO.:			JP 1988-43440	19880226 <

AB The title photog. material contains a shift coupler in which the wavelength of the absorption maximum of the dye released is shorter in the bonded state before release in comparison with that in the free state and has hydrophilic

colloidal layers hardened by a hardening agent through activation of carboxylic groups.

IT 125329-20-6

RL: USES (Uses)

(photog. yellow shift coupler)

RN 125329-20-6 CAPLUS

CN Benzothiazolium,

chloro-5-[(dodecyloxy)carbonyl]phenyl]amino]carbonyl]-2-(4-methoxyphenyl)2-oxoethoxy]phenyl]-3-methyl-, iodide (9CI) (CA INDEX NAME)

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L87 ANSWER 54 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1990:431810 CAPLUS Full-text

DOCUMENT NUMBER: 113:31810
ORIGINAL REFERENCE NO.: 113:5319a,5322a

TITLE: Silver halide color photographic material containing

azole compound as cyan coupler

INVENTOR(S): Fukunaga, Hiroo; Yamakawa, Kazuyoshi; Furusawa,

Genichi

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 44 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01250949	A	19891005	JP 1988-76403	19880331 <
PRIORITY APPLN. INFO.:			JP 1988-76403	19880331 <
OTHER SOURCE(S):	MARPAI	113:31810		
GI				

AB A Ag halide color photog. material having excellent color reproducibility contains an azole compound [I; R = H, blocking group; Z, Z1, Z2 = (substituted) methine, N, but the substituent \neq OH, acyloxy, sulfonyloxy] as a cyan coupler. A color photog. film having II as a cyan coupler was processed to give images showing excellent color reproducibility and colorfastness on storage at 60° and 70% relative humidity.

IT 127828-92-6

RL: TEM (Technical or engineered material use); USES (Uses) (cyan photog. coupler)

RN 127828-92-6 CAPLUS

CN Tetradecanamide, 2-[2-chloro-4-[(3-chloro-4-hydroxyphenyl)sulfonyl]phenoxy]-N-[2-(3-chloro-5-ethyl-1H-pyrazol-4-yl)phenyl]- (CA INDEX NAME)

$$CI$$
 NH
 CI
 $CH_2)_{11}$
 Me
 CI
 OH
 CI
 OH

ACCESSION NUMBER: 1990:188876 CAPLUS Full-text

DOCUMENT NUMBER: 112:188876

ORIGINAL REFERENCE NO.: 112:31745a,31748a

TITLE: High-sensitivity color photographic material with

improved image sharpness and shelf life

INVENTOR(S): Hirabayashi, Shigeto PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 14 pp.

CODEN: JKXXAF

DOCUMENT TYPE: %atent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01191142	A	19890801	JP 1988-15453	19880126 <
PRIORITY APPLN. INFO.:			JP 1988-15453	19880126 <

AB In the title photog. material, ≥1 of the Ag halide emulsion layers contains a benzoylacetanilide derivative yellow photog. coupler and another yellow photog. coupler which, upon reaction with an oxidized aromatic primary amine color developer, releases a dye whose absorption maximum is at a shorter wavelength prior to its release.

IT 125329-20-6

RL: USES (Uses)

(yellow photog. coupler)

RN 125329-20-6 CAPLUS

CN Benzothiazolium,

Me— (CH₂)₁₁-0-C

NH

C1

OMO

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L87 ANSWER 56 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1990:188873 CAPLUS Full-text

DOCUMENT NUMBER: 112:188873

ORIGINAL REFERENCE NO.: 112:31745a,31748a

TITLE: Silver halide photographic material with improved

sharpness

INVENTOR(S): Yagi, Toshihiko; Nakagawa, Satoshi

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 22 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01188847	A	19890728	JP 1988-13929	19880125 <
PRIORITY APPLN. INFO.:			JP 1988-13929	19880125 <

AB In a Ag halide photog, material containing ≥1 Ag halide emulsion layer containing a coupler, ≥1 of the emulsion layers contains Ag halide grains containing ≥2 phases differing in AgI content with an. AgI content 2-20 mol% and the average AgI content higher than that in the peripheral regions and a coupler which contains a dye- or dye precursor-releasing group bonded directly or via a timing group to the active site with the above dye showing a shorter absorption maximum wave length before its release.

IT 125329-20-6

RL: TEM (Technical or engineered material use); USES (Uses) (photog. coupler)

RN 125329-20-6 CAPLUS

CN Benzothiazolium,

chloro-5-[(dodecyloxy)carbonyl]phenyl]amino]carbonyl]-2-(4-methoxyphenyl)-2-oxoethoxy]phenyl]-3-methyl-, iodide (9CI) (CA INDEX NAME)

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L87 ANSWER 57 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1990:174016 CAPLUS Full-text

DOCUMENT NUMBER: 112:174016

ORIGINAL REFERENCE NO.: 112:29279a,29282a

TITLE: Stable pesticidal emulsions

AUTHOR(S): Curtis, Ralston

CORPORATE SOURCE: Sandoz Crop Prot., USA

SOURCE: Research Disclosure (1989), 308, 981-2 (No. 308103)

CODEN: RSDSBB; ISSN: 0374-4353

DOCUMENT TYPE: Journal; Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
RD 308103		19891210	RD 1989-308103	19891210 <
PRIORITY APPLN. INFO.:			RD 1989-308103	19891210 <

AB Stable emulsifiable concs. containing 1-20% by weight of a difficult-to-dissolve crystalline pesticide, 45-90% by weight of 1-octyl-2-pyrrolidinone, 5-30% by weight of 1-methyl-2-pyrrolidinone, cyclohexanone, or α -butyrolactone, 10-50% by weight of methylated

naphthalene or a xylene-range aromatic solvent or a mixture of 2 or more xylene-range aromatic solvents and 1-10% by weight of a surfactant are prepared Pesticides prepared from these concs. were effective for controlling beet armyworm (Spodoptera exigua) on lima bean plants.

IT 112737-76-5 126353-95-5

RL: BIOL (Biological study)

(stable pesticidal emulsion containing)

RN 112737-76-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 126353-95-5 CAPLUS

CN Benzamide,

2-chloro-N-[[[2,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F3C \longrightarrow \begin{array}{c} C1 \\ NH - C - NH - C \\ C1 \\ \end{array}$$

L87 ANSWER 58 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1990:88181 CAPLUS Full-text

DOCUMENT NUMBER: 112:88181

ORIGINAL REFERENCE NO.: 112:14831a,14834a

TITLE: Fog-resistant silver halide photographic material

INVENTOR(S): Sakamoto, Hidekazu PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 18 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01196045	A	19890807	JP 1988-20871	19880130 <
PRIORITY APPLN. INFO.:			JP 1988-20871	19880130 <

GΙ

AB The material contains I (Z = group forming C-, N-, and/or S-containing 5- or 6-membered heterocycle; M = H, alkali metal, NH4, protective group) and ≥1 coupler bound with a dye or its precursor, released by the reaction with an oxidant of aromatic primary amine coloring developer, directly or via timing group at the activation position, and the absorption maximum of the dye or its precursor shows blue shift before it is released. A Ag halide photog. emulsion containing II and III was applied on a cellulose triacetate film support to give a color photog. material which showed high durability and fog resistance.

IT 125329-20-6

RL: TEM (Technical or engineered material use); USES (Uses) (photog. yellow coupler)

RN 125329-20-6 CAPLUS

CN Benzothiazolium,

Me— (CH₂)₁₁— O— C

Me

NH

C1

OMe

OMe

OMe

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L87 ANSWER 59 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1990:55721 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 112:55721
ORIGINAL REFERENCE NO.: 112:9571a,9574a

TITLE: Synthesis and antibacterial activity of some new

selenadiazoles and thiadizoles containing amino acid

moieties

AUTHOR(S): Bayoumy, B. E.; Deeb, A.; El-Mobayed, M.; Abd-Alla, M.

Α.

CORPORATE SOURCE: Fac. Sci., Zagazig Univ., Zagazig, Egypt

SOURCE: Egyptian Journal of Chemistry (1988), Volume Date

1987, 30(1), 53-61

CODEN: EGJCA3; ISSN: 0367-0422

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 112:55721

GI

- AB Title compds. I (R = piperazino, morpholino, piperidino, 2-pyridylamino, ethylamino, etc.; Z = Se, S) were prepared by treating semicarbazones II with SeO2/AcOH or SOC12 resp. Bactericidal and fungicidal activity of I were determined
- IT 111281-88-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation, bactericidal, and fungicidal activity of)

RN 111281-88-0 CAPLUS

CN Acetamide, 2-[(4-chlorophenyl)amino]-N-[4-(1,2,3-thiadiazol-4-yl)phenyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

L87 ANSWER 60 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1990:14188 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 112:14188

ORIGINAL REFERENCE NO.: 112:2443a,2446a

TITLE: Silver halide color photographic material containing

cyan coupler

INVENTOR(S): Tachibana, Kimie; Kaneko, Yutaka

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 13 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01028638	A	19890131	JP 1987-184554	19870723 <
PRIORITY APPLN. INFO.:			JP 1987-184554	19870723 <
GI				

In the title photog. material, ≥ 1 of red-sensitive Ag halide emulsion layer contains a coupler I [R1, R2 = substituent; Y, R3 = H, substituent; X = H, group to be released upon reaction with an oxidized developer]. The coupler is a pyrazoloimidazole cyan coupler. Cyan images show excellent spectral characteristics, and heat—and moisture—resistance. II was an example of T.

IT 124171-49-9

RL: USES (Uses)

(cyan coupler, color photog. material containing)

RN 124171-49-9 CAPLUS

CN 1H-Imidazo[1,2-b]pyrazole-6-carboxamide,

2-[4-[[2-(2-chlorophenoxy)-1-oxotetradecyl]amino]phenyl]-3-cyano-N-ethyl-(CA INDEX NAME)

IT 124171-39-7P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and use of, as cyan coupler)

RN 124171-39-7 CAPLUS

CN 1H-Imidazo[1,2-b]pyrazole-6-carboxamide,

7-chloro-2-[4-[[2-(2-chlorophenoxy)-1-oxotetradecyl]amino]phenyl]-N-ethyl-3-methyl- (CA INDEX NAME)

L87 ANSWER 61 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1989:574087 CAPLUS Full-text

DOCUMENT NUMBER: 111:174087

ORIGINAL REFERENCE NO.: 111:29006h,29007a TITLE: Preparation of

N-benzoyl-N'-(4-pyrazol-1-yl-phenyl)ureas as

pesticides

INVENTOR(S): Neubauer, Hans Juergen; Kuenast, Christoph;

Hofmeister, Peter

PATENT ASSIGNEE(S): BASF A.-G., Fed. Rep. Ger.

SOURCE: Ger. Offen., 14 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

DE 3732541 A1 19890413 DE 1987-3732541 19870926 <-PRIORITY APPLN. INFO.: DE 1987-3732541 19870926 <-OTHER SOURCE(S): CASREACT 111:174087; MARPAT 111:174087

GΙ

The title compds. (I; R1, R3, R4 = H, halo; R2 = halo; R5-R7 = R1, C1-8 alkyl, C1-4 haloalkyl) were prepared 4-(O2N)C6H4NHNH2.HCl was refluxed 1 h with (CF3CO)2CH2 in EtOH and the product mixture stirred 8 h with MeSO2Cl in CH2Cl2 containing Et3N to give phenylpyrazole II (R = NO2) which was heated 2 h at 60° with ZnCl2 and HCl in EtOH to give II (R = NH2). The latter was stirred 2 h at 40° with 2,6-F2C6H3CONCO in THF to give II (R = 2,6-F2C6H3CONHCONH) which was lethal to larvae of Aedes aegypti at 0.002 ppm.

ΙT 112737-04-99 112737-05-02 112737-21-02 112737-22-19 112737-62-9P 112762-62-6P 123066-42-29 123066-43-3P 123066-44-4P 123066-45-5P 123066-46-69 123066-47-79 123066-48-8P 123066-49-99 123066-50-2P 123066-51-3P 123066-52-4P 123066-53-5P 123066-55-7P 123066-56-8P 123066-54-6P 123066-57-9P 123066-58-0P 123066-59-1P 123066-62-6P 123066-60-4P 123066-61-5P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as pesticide)

RN 112737-04-9 CAPLUS

CN Benzamide, N-[[[4-(4-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 112737-05-0 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-(1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-21-0 CAPLUS

CN Benzamide, N-[[[4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-22-1 CAPLUS

CN Benzamide, N-[[[3-chloro-4-(4-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & &$$

RN 112737-62-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112762-62-6 CAPLUS

CN Benzamide, N-[[[4-[3,5-bis(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 123066-42-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-(1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

RN 123066-43-3 CAPLUS

CN Benzamide, 2,6-difluoro-N-[[[4-(1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 123066-44-4 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-(4-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 123066-45-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-(4-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 123066-46-6 CAPLUS

CN Benzamide, N-[[[4-[3,5-bis(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

RN 123066-47-7 CAPLUS

CN Benzamide,

2,6-difluoro-N-[[[4-[5-methyl-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 123066-48-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[5-methyl-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} F3C & N & O & O \\ \hline & NH & C & NH & C \\ \hline & NH & C & O \\ \hline \end{array}$$

RN 123066-49-9 CAPLUS

CN Benzamide,

2,6-difluoro-N-[[[4-[3-methyl-5-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} \text{Me} & \bigcirc & \bigcirc & \bigcirc & \bigcirc & \bigcirc & \\ & & & \text{NH-C-NH-C} \\ & & & & \text{NH-C-NH-C} \\ & & & & & \\ \end{array}$$

RN 123066-50-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[3-methyl-5-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 123066-51-3 CAPLUS

CN Benzamide,

N-[[[4-(3,5-dimethyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c} \text{Me} \\ \\ \text{Me} \\ \end{array}$$

RN 123066-52-4 CAPLUS

CN Benzamide, 2,6-dichloro-N-[[[4-(3,5-dimethyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 123066-53-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-(3,5-dimethyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 123066-54-6 CAPLUS

CN Benzamide, N-[[[3-chloro-4-(3-methyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 123066-55-7 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-(3-methyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 123066-56-8 CAPLUS

CN Benzamide, 2,6-difluoro-N-[[[4-(3-methyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 123066-57-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-(3-methyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} Me & & N \\ \hline & N \\ \hline & NH \\ \hline & C \\ NH \\ \hline & C \\ \hline & C \\ \end{array}$$

RN 123066-58-0 CAPLUS

CN Benzamide, N-[[[4-(5-chloro-3-methyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} \text{Me} & \\ \hline \\ \text{NH} & \\ \hline \\ \text{NH} & \\ \hline \\ \text{NH} & \\ \hline \\ \text{F} \\ \hline \end{array}$$

RN 123066-59-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-(5-chloro-3-methyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 123066-60-4 CAPLUS

CN Benzamide,

N-[[[4-(3-ethyl-4-methyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} \text{Et} & \text{N} & \text{N} \\ \text{Me} & \text{NH} & \text{C} & \text{NH} \\ \end{array}$$

RN 123066-61-5 CAPLUS

CN Benzamide, 2,6-dichloro-N-[[[4-(3-ethyl-4-methyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 123066-62-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-(3-ethyl-4-methyl-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS)

L87 ANSWER 62 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1988:619497 CAPLUS Full-text

DOCUMENT NUMBER: 109:219497

ORIGINAL REFERENCE NO.: 109:36159a,36162a

TITLE: Silver halide color photographic light-sensitive

material containing pyrazoloazole type cyan coupler

INVENTOR(S): Tachibana, Kimie; Kaneko, Yutaka; Ishii, Fumio

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Eur. Pat. Appl., 168 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 269436	A2	19880601	EP 1987-310417	19871125 <

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EP 269436
                         A3
                               19880914
    EP 269436
                         В1
                               19910306
    EP 269436
                         B2
                               19941102
        R: DE, FR, GB, IT, NL
                                          JP 1987-294701
    JP 64000554
                        Α
                              19890105
                                                                19871121 <--
    JP 2517334
                        В2
                               19960724
    JP 64000555
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                              19890105
                                          JP 1987-294702
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    US 4873183
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                              19891010
                                          US 1988-291351
                                                                19881229 <--
PRIORITY APPLN. INFO .:
                                          JP 1986-280164
                                                             A 19861125 <--
                                          JP 1986-313455
                                                             A 19861227
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                                           JP 1987-47323
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                                          JP 1987-62162
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                                           JP 1987-184552
                                                              A 19870723
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                                           JP 1986-313458
                                                              A1 19861227 <--
                                           US 1987-124987
                                                              A1 19871124 <--
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GΙ For diagram(s), see printed CA Issue.

AΒ A light-sensitive Ag halide color photog. material comprises a support, a red light-sensitive Ag halide emulsion layer containing a pyrazoloazole-type cyan dye-forming coupler having ≥1 electron-attracting group in a substitutable position except the active site of the coupler. The cyan dye-forming coupler is I [R1 = H, substituent; R2 = H, substituent provided that ≥ 1 of R1 and R2 (≥ 1 of R2 when there are more than 2 R2 s) is electron-attracting group; Z = group of nonmetal atoms necessary to complete a N-containing heterocyclic ring; R2 being connected with a C atom of the heterocyclic ring; X = H, substituent capable of being split off upon reaction with the oxidized product of a color developing agent; n = 1, 2]. The absorption maximum of the cyan dye-forming coupler is 580-710 nm. The red light-sensitive emulsion layer may contain a spectral sensitizing dye. A color photog. material containing the coupler provides excellent color reproduction

ΙT 117507-99-0 117543-05-2

RL: TEM (Technical or engineered material use); USES (Uses) (cyan photog. coupler)

117507-87-6 CAPLUS

Tetradecanamide, CN

2-(2-chlorophenoxy)-N-[4-[6-(2,3,4,5,6-pentafluorophenyl)-1H-pyrazolo[5,1-c]-1,2,4-triazol-3-yl]phenyl]- (CA INDEX NAME)

RN117507-99-0 CAPLUS

Tetradecanamide,

2-(2-chlorophenoxy)-N-[4-[7-chloro-6-(trifluoromethyl)-3Hpyrazolo[1,5-b][1,2,4]triazol-2-yl]phenyl]- (CA INDEX NAME)

117543-05-2 CAPLUS RN

Tetradecanamide, 2-[2-chloro-4-[(3-chloro-4-CN $\label{lem:hydroxyphenyl} \verb| hydroxyphenyl| \verb| sulfonyl| -N-[3-[3-methyl-5-(phenylsulfonyl)-1H-1] - (phenylsulfonyl) -1H-1 - (phenylsulfonyl) -1$ pyrazolo[1,5-b]pyrazol-2-yl]phenyl]- (CA INDEX NAME)

OS.CITING REF COUNT: THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)

L87 ANSWER 63 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1988:221637 CAPLUS Full-text

DOCUMENT NUMBER: 108:221637 ORIGINAL REFERENCE NO.: 108:36383a,36386a

TITLE: Synthesis and antibacterial testing of some new

selenadiazole and thiadiazole containing amino acid

moieties

AUTHOR(S): Bayoumi, B. E.; Abd-Alla, M. A.; Ahmed, A. N.

CORPORATE SOURCE: Fac. Sci., Zagazig Univ., Assiut, Egypt

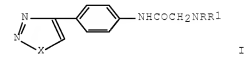
SOURCE: Bulletin of Pharmaceutical Sciences, Assiut University

(1986), 9(2), 66-77

CODEN: BPAUEC; ISSN: 1110-0052

DOCUMENT TYPE: Journal LANGUAGE: English

GΙ



An umber of glycylaminophenylselenadiazoles and -thiadiazoles (e.g., I; RR1N = morpholino, piperidino, EtNH; X = Se, S) were prepared and evaluated for antimicrobial activity. Thus, 4-MeCOC6H4NHCOCH2Cl was treated with RR1NH to give 4-MeCOC6H4NHCOCH2NRR1. Treating the latter compds. with H2NCONHNH2 gave 4-H2NCONHN:CMeC6H4NHCOCH2NRR1, which underwent oxidative cyclization with SeO2 in AcOH or SOCl2 to give I. The min. inhibitory concentration for I (RR1N = 2-thiazolylamino; X = Se, S) against several bacteria including Staphylococcus aureus, Klebsiella pneumoniae, and Escherichia coli were lower than those for tetracycline.

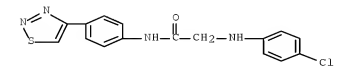
IT 111281-88-0P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and antimicrobial activity of)

RN 111281-88-0 CAPLUS

CN Acetamide, 2-[(4-chlorophenyl)amino]-N-[4-(1,2,3-thiadiazol-4-yl)phenyl]- (CA INDEX NAME)



OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)

L87 ANSWER 64 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1988:94547 CAPLUS <u>Full-text</u> DOCUMENT NUMBER: 108:94547

ORIGINAL REFERENCE NO.: 108:15555a,15558a

TITLE: Preparation of N-benzoyl-N'-(heterocyclylphenyl)ureas

as insecticides and acaricides

INVENTOR(S): Carney, Robert L.; Gruber, John M.; Lui, Alfred S.

PATENT ASSIGNEE(S): Sandoz A.-G., Switz.; Sandoz-Patent-G.m.b.H.;

Sandoz-Erfindungen Verwaltungsgesellschaft m.b.H.

SOURCE: Eur. Pat. Appl., 26 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
EP 242322	A2	19871021	EP 1987-810137		19870311 <
EP 242322	A3	19890322			
EP 242322	B1	19921216			
R: AT, BE, CH,	DE, ES	FR, GB,	GR, IT, LI, NL		
AT 83477	T	19930115	AT 1987-810137		19870311 <
ES 2053579	Т3	19940801	ES 1987-810137		19870311 <
DK 8701345	A	19870919	DK 1987-1345		19870316 <
AU 8770048	A	19870924	AU 1987-70048		19870316 <
AU 602884	B2	19901101			
IL 81900	A	19910610	IL 1987-81900		19870316 <
CN 87102145	A	19870930	CN 1987-102145		19870317 <
CN 1017397	В	19920715			
JP 62230765	A	19871009	JP 1987-65533		19870317 <
BR 8701205	A	19880112	BR 1987-1205		19870317 <
HU 43941	A2	19880128	HU 1987-1157		19870317 <
HU 203945	В	19911128			
SU 1491333	A3	19890630	SU 1987-4202182		19870317 <
ZA 8701996	A	19881026	ZA 1987-1996		19870318 <
PRIORITY APPLN. INFO.:			US 1986-840814	A	19860318 <
			EP 1987-810137	A	19870311

$$\begin{array}{c} X1 \\ X2 \\ X3 \\ X2 \end{array}$$

The title compds. [I; A = N, R4C; B = N, R3C; R1, R4 = H, halo, C1-8 haloalkyl, C1-8 (halo)alkoxy, C1-8 (halo)alkylthio, (un)substituted aryl, aryloxy, arylthio; R2, R3 = H, halo, cyano, C02R5, C1-8 (halo)alkyl, C1-8 (halo)alkoxy, C1-8 (halo)alkylthio, (un)substituted aryl, aryloxy, arylthio; R1R2, R2R3 = (un)substituted, (un)saturated C4 bridging group; R5

= H, C1-8 alkyl; X1-X3, X5 = H, halo, C1-4 alkyl; X4 = H, halo, C1-4 (halo)alkyl, C02R5; X6 = H, halo, C1-8 alkyl, C02R5; Y = 0, S] were prepared as chitin inhibitors, useful as acaricides and insecticides. 2,6-F2C6H3CONCO was added dropwise to a solution of p-(4-chloropyrazol-1-yl)aniline in CH2Cl2 and the mixture stirred 30 min at room temperature to give I (A = N, B = CH, R1 = X3-X6 = H, R2 = Cl, X1 = X2 = F, Y = 0). I showed insecticidal activity when applied topically at 0.004-0.070 μ g/insect to third instar larvae of Heliothis virescens.

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ΙT
     112736-96-62
                    112736-97-7P
                                    112736-98-8P
     112736-99-9P
                    112737-00-5P
                                    112737-01-6P
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     RL: AGR (Agricultural use); BAC (Biological activity or effector, except
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RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as insecticide and acaricide)

RN 112736-96-6 CAPLUS

CN

Benzamide, N-[[[3,5-dichloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112736-97-7 CAPLUS

CN Benzamide, N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-3,5-dimethylphenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\$$

RN 112736-98-8 CAPLUS

CN Benzamide, N-[[[4-(4-bromo-1H-pyrazol-1-y1)-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

RN 112736-99-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3,4-dibromo-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-00-5 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F3C \longrightarrow N \\ C1 \\ C1 \\ C1 \\ NH-C \\ NH-$$

RN 112737-01-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-02-7 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-03-8 CAPLUS

CN Benzamide, N-[[[4-[3-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

CN Benzamide, N-[[[4-(4-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 112737-05-0 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-(1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 112737-06-1 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(4-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-07-2 CAPLUS

CN Benzamide, N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

CN Benzamide, N-[[[4-[3,4-bis(trifluoromethyl)-1H-pyrazol-1-y1]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F3C$$

$$NH-C-NH-C$$

$$F3C$$

RN 112737-09-4 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$C1 \longrightarrow NH - C - NH - C \longrightarrow F$$

RN 112737-10-7 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3,4-dibromo-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-11-8 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array}$$

RN 112737-12-9 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

PAGE 1-A



- RN 112737-13-0 CAPLUS
- CN Benzamide, N-[[[3,5-dichloro-4-[3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

- RN 112737-14-1 CAPLUS
- CN Benzamide, N-[[[3,5-dichloro-4-[4-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-15-2 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F3C \longrightarrow N \longrightarrow NH \longrightarrow C \longrightarrow NH \longrightarrow C \longrightarrow F$$

RN 112737-16-3 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(1,1-dimethylethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-17-4 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3,4-dichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$C1 \longrightarrow NH - C - NH - C \longrightarrow F$$

RN 112737-18-5 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-[4-chloro-3-(trifluoromethyl)-1H-pyrazol-1-

yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F_{3}C$$

$$NH-C-NH-C$$

$$F$$

RN 112737-19-6 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F3C \longrightarrow NH-C-NH-C-NH-C$$

RN 112737-20-9 CAPLUS

CN Benzamide, N-[[[4-[3,5-bis(trifluoromethyl)-1H-pyrazol-1-y1]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F_3C \xrightarrow{\qquad \qquad N \\ CF_3} C1 \xrightarrow{\qquad \qquad NH-C-NH-C} F$$

RN 112737-21-0 CAPLUS

CN Benzamide, N-[[[4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-22-1 CAPLUS

CN Benzamide, N-[[[3-chloro-4-(4-chloro-1H-pyrazol-1-

yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 112737-23-2 CAPLUS

CN Benzamide, N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-y1]-3-(trifluoromethyl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ &$$

RN 112737-24-3 CAPLUS

CN Benzamide, N-[[[4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]-3-(trifluoromethyl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F3C \longrightarrow NH-C-NH-C$$

RN 112737-25-4 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-26-5 CAPLUS

CN Benzamide, N-[[[4-[4-bromo-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-27-6 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3,4,5-tribromo-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-28-7 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-[3-chloro-5-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$C1 \xrightarrow{N} NH C NH C F$$

RN 112737-29-8 CAPLUS

CN Benzamide, N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-2-fluorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-30-1 CAPLUS

CN Benzamide,

 $\begin{tabular}{ll} $N-[[[5-chloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-2-fluorophenyl]amino]carbonyl]-2,6-difluoro-(CA INDEX NAME) \\ \end{tabular}$

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 112737-31-2 CAPLUS

CN Benzamide, N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-3-methylphenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

RN 112737-32-3 CAPLUS

CN Benzamide,

N-[[[5-chloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-2-methylphenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-33-4 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4-(2,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

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RN 112737-34-5 CAPLUS

CN Benzamide, N-[[[4-[4-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

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RN 112737-35-6 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4-chloro-3-(4-chlorophenyl)-5-methoxy-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-36-7 CAPLUS

CN 1H-Pyrazole-4-carboxylic acid, 1-[2,6-dichloro-4-[[[(2,6-difluorobenzoyl)amino]carbonyl]amino]phenyl]-, methyl ester (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 112737-37-8 CAPLUS

CN Benzoic acid, 2-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-5-[[[(2,6-difluorobenzoyl)amino]carbonyl]amino]-, methyl ester (CA INDEX NAME)

RN 112737-38-9 CAPLUS

CN Benzamide, N-[[[3-chloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$F3C \longrightarrow NH-C-NH-C \longrightarrow F$$

RN 112737-39-0 CAPLUS

CN Benzamide,

N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3-chlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & &$$

CN Benzamide, N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3-(trifluoromethyl)phenyl]amino]carbonyl]-2,6-difluoro-(CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 112737-41-4 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(2,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-42-5 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(3,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-43-6 CAPLUS

CN Benzamide, N-[[[4-[3-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-44-7 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(4-fluorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\underbrace{ \left(\begin{array}{c} C \\ F \end{array} \right) \left(\begin{array}{c} C \\ V \end{array} \right) \left(\begin{array}{c}$$

RN 112737-45-8 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-[3-[3-(trifluoromethyl)phenyl]-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-46-9 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(5-chloro-2-thienyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ &$$

RN 112737-47-0 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(2-naphthalenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & &$$

RN 112737-48-1 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(4-cyano-1H-pyrazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-49-2 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-50-5 CAPLUS

CN Benzamide, N-[[[3-chloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

$$F3C \longrightarrow NH-C-NH-C$$

RN 112737-51-6 CAPLUS

CN Benzamide,

N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3-chlorophenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-52-7 CAPLUS

CN Benzamide, N-[[[4-(4-bromo-1H-pyrazol-1-y1)-3- (trifluoromethyl)phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

RN 112737-53-8 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(2,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-54-9 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(3,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

CN Benzamide, N-[[[3,5-dichloro-4-[3-(4-fluorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

RN 112737-56-1 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-[3-[3-(trifluoromethyl)phenyl]-lH-pyrazol-l-yl]phenyl]amino]carbonyl]-2-fluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &$$

RN 112737-59-4 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-5-fluoro- (CA INDEX NAME)

RN 112737-60-7 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-6-fluoro- (CA INDEX NAME)

RN 112737-61-8 CAPLUS

CN Benzamide, 2,6-dichloro-N-[[[3,5-dichloro-4-[4,5-dichloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F_{3}C$$

$$C_{1}$$

$$C_{2}$$

$$C_{1}$$

$$C_{1}$$

$$C_{2}$$

$$C_{3}$$

$$C_{4}$$

$$C_{1}$$

$$C_{2}$$

$$C_{3}$$

$$C_{4}$$

$$C_{5}$$

$$C_{6}$$

$$C_{7}$$

$$C_{1}$$

$$C_{7}$$

$$C_{8}$$

$$C_{1}$$

$$C_{1}$$

$$C_{1}$$

$$C_{1}$$

$$C_{2}$$

$$C_{3}$$

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$$C_{8}$$

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$$C_{1}$$

$$C_{1}$$

$$C_{2}$$

$$C_{3}$$

$$C_{4}$$

$$C_{5}$$

$$C_{7}$$

$$C_{7}$$

$$C_{8}$$

$$C_{$$

RN 112737-62-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-63-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(4-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-64-1 CAPLUS

CN Benzamide, N-[[[4-[3,4-bis(trifluoromethyl)-1H-pyrazol-1-y1]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

$$F_{3}C$$

$$NH-C-NH-C$$

$$C_{1}$$

$$C_{2}$$

$$C_{1}$$

$$C_{1}$$

$$C_{2}$$

$$C_{3}$$

$$C_{4}$$

$$C_{2}$$

$$C_{3}$$

$$C_{4}$$

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RN 112737-65-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3-chloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$C1 \longrightarrow NH - C - NH - C$$

RN 112737-66-3 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[4-(4-chlorophenyl)-lH-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

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RN 112737-67-4 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[4-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c}
C1 & C1 \\
NH-C-NH-C
\end{array}$$

RN 112737-68-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F_3C$$

$$\begin{array}{c} C1 \\ NH-C-NH-C \\ \hline \\ C_1 \end{array}$$

RN 112737-69-6 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[3-(1,1-dimethylethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-70-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[4-chloro-3-(4-chlorophenyl)-1H-

pyrazol-1-y1]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-71-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[4-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F3C$$

$$C1$$

$$NH-C-NH-C$$

RN 112737-72-1 CAPLUS

CN Benzamide, N-[[[4-[3,5-bis(trifluoromethyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

$$F_3C \xrightarrow{C1} NH - C - NH - C$$

$$CT - NH - C$$

RN 112737-73-2 CAPLUS

CN Benzamide, N-[[[4-[4-bromo-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & & & & & & & & & & & & & \\ & & & & & & & & & & & & & & \\ & & & & & & & & & & & & \\ & & & & & & & & & & & \\ & & & & & & & & & & \\ & & & & & & & & & \\ & & & & & & & & & \\ & & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & \\ & & \\ &$$

RN 112737-74-3 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3,4,5-tribromo-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-75-4 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-chloro-5-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-76-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3,4,5-trichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-77-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-3,5-dimethylphenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-79-8 CAPLUS
CN Benzamide, N-[[[4-(4-bromo-1H-pyrazol-1-yl)-3(trifluoromethyl)phenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

$$\begin{array}{c} C1 \\ NH-C-NH-C \\ \end{array}$$

RN 112737-80-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-(2-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-81-2 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[3-(2,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$0 = \begin{bmatrix} C & & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & \\ & & \\ & & \\ &$$

RN 112737-82-3 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[3-(3,4-dichlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-83-4 CAPLUS

CN Benzamide, N-[[[4-[3-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

RN 112737-84-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-5-methylphenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-85-6 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-(4-fluorophenyl)-1H-pyrazol-1-

yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

RN 112737-86-7 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[3-[3-(trifluoromethyl)phenyl]-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-87-8 CAPLUS

CN Benzamide,

2-chloro-N-[[[3,5-dichloro-4-[3-(5-chloro-2-thienyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

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RN 112737-88-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-(2-naphthalenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-89-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-[5-chloro-3-(trifluoromethyl)-1H-

pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112737-90-3 CAPLUS

CN Benzamide, N-[[[4-[5-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

RN 112737-91-4 CAPLUS

CN Benzamide, N-[[[4-[5-(4-bromophenyl)-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-92-5 CAPLUS

CN Benzamide, N-[[[4-[5-(4-bromophenyl)-4-chloro-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

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RN 112737-93-6 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(4-methylphenyl)-1H-pyrazol-1-

yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & &$$

RN 112737-94-7 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-[3-(4-methylphenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

RN 112737-96-9 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(3-chloro-1H-indazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-97-0 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(5,6-dichloro-1H-benzimidazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c} C1 \\ C1 \\ \end{array}$$

RN 112737-98-1 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(5-chloro-1H-benzotriazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112737-99-2 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(6-chloro-1H-benzotriazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112738-00-8 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-(5,6-dichloro-1H-benzotriazol-1-yl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112738-01-9 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(5,6-dichloro-1H-benzimidazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112738-02-0 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-(5,6-dichloro-1H-benzimidazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112738-03-1 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(5-chloro-1H-benzotriazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112738-04-2 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(5,6-dichloro-1H-benzotriazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112738-05-3 CAPLUS

CN Benzamide, N-[[[3-chloro-4-[5,6-dichloro-2-(trifluoromethyl)-1H-benzimidazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} C1 & & \\ \hline \\ C1 & & \\ \hline \\ C1 & & \\ \hline \\ CF3 & \\ \hline \\ NH- \\ \hline \\ NH- \\ \hline \\ F & \\ \hline \\ F & \\ \hline \\ \end{array}$$

RN 112738-06-4 CAPLUS

CN Benzamide,

N-[[[3,5-dichloro-4-(2H-indazol-2-y1)pheny1]amino]carbony1]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

RN 112738-07-5 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(2H-indazol-2-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112762-62-6 CAPLUS

CN Benzamide, N-[[[4-[3,5-bis(trifluoromethyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

RN 112762-63-7 CAPLUS

CN Benzamide, 2-chloro-N-[[[3,5-dichloro-4-(3,4-dichloro-1H-pyrazol-1-yl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$\begin{array}{c|c} C1 & NH-C-NH-C \\ \hline \end{array}$$

RN 112762-64-8 CAPLUS

CN Benzamide, 2-chloro-N-[[[3-chloro-4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112762-65-9 CAPLUS

CN Benzamide,

2-chloro-N-[[[4-[4-chloro-3-(4-chlorophenyl)-1H-pyrazol-1-yl]-3-(trifluoromethyl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 112762-66-0 CAPLUS

CN Benzamide,

2-chloro-N-[[[4-[5-chloro-3-(trifluoromethyl)-1H-pyrazol-1-yl]-3-(trifluoromethyl)phenyl]amino]carbonyl]- (CA INDEX NAME)

$$F_{3}C$$

$$NH-C-NH-C$$

$$C1$$

$$C1$$

$$C_{F3}$$

RN 112762-67-1 CAPLUS

CN Benzamide, N-[[[4-[5-(4-bromophenyl)-4-chloro-1H-pyrazol-1-yl]-3,5-dichlorophenyl]amino]carbonyl]-2-chloro- (CA INDEX NAME)

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RN 112785-82-7 CAPLUS

CN Benzamide, N-[[[3,5-dichloro-4-[3-(2-chlorophenyl)-1H-pyrazol-1-yl]phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\bigcap_{F}^{C} \bigcap_{C-NH-C-NH}^{C-NH-C-NH} \bigcap_{C1}^{C1} \bigcap_{N}^{C1} \bigcap_{N$$

OS.CITING REF COUNT: 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS)

L87 ANSWER 65 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1987:617551 CAPLUS Full-text

DOCUMENT NUMBER: 107:217551

ORIGINAL REFERENCE NO.: 107:34903a,34906a

TITLE: Synthesis and antibacterial activity of some new

selenadiazole- and thiadiazole-containing amino acid

moieties

AUTHOR(S): Bayoumy, B. E.; Deeb, A.; El-Mobayed, M.; Abd-Alla, M.

Α.

I

CORPORATE SOURCE: Fac. Sci., Zagazig Univ., Egypt

SOURCE: Egyptian Journal of Pharmaceutical Sciences (1986),

27(1-4), 17-26

CODEN: EJPSBZ; ISSN: 0301-5068

DOCUMENT TYPE: Journal LANGUAGE: English

GΙ

NHCOCH2R

AB Title compds. I (X = Se, S; R = piperidino, piperazino, morpholino, NHR1; R1 = Et, 2-pyridyl, 2-thiazolyl, C6H4OMe-p, C6H4Cl-p), which were prepared, showed bactericidal and fungicidal activity. Heating 4-RCH2CONHC6H4CMe:NNHCONH2 (same R) with SeO2 yielded I (X = Se).

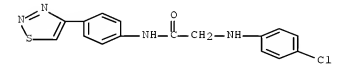
IT 111281-88-0P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and fungicidal activity of)

RN 111281-88-0 CAPLUS

CN Acetamide, 2-[(4-chlorophenyl)amino]-N-[4-(1,2,3-thiadiazol-4-yl)phenyl](CA INDEX NAME)



OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

L87 ANSWER 66 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1987:449424 CAPLUS Full-text

DOCUMENT NUMBER: 107:49424

ORIGINAL REFERENCE NO.: 107:8055a,8058a

TITLE: Silver halide color photographic couplers

INVENTOR(S): Yamada, Kozaburo; Ichijima, Yasushi; Obayashi, Keiji

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 33 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
JP 61233741	A	19861018	JP 1985-72378	19850405 <	
JP 06080458	В	19941012			
PRIORITY APPLN. INFO.:			JP 1985-72378	19850405 <	

GI For diagram(s), see printed CA Issue.

AB A Ag halide color photog. material contains ≥1 coupler selected from the compds. having the general formulas of I, II, III, IV, V, and VI [A = coupler group which splits off from O upon reaction with an oxidized developer; PUG = photog. useful group; R = H, substituent; R1 = acylamino, sulfonamido, ureido, sulfamoylamino, amino; Z = organic moiety fused to the benzene ring to form a 5-6-membered ring other than a benzene ring; 1, m, n = 1, 2; n + m ≤3 in I; n + m + 1 ≤4 in II].

IT 109117-35-3P

RL: PREP (Preparation)

(preparation of, as photog. development inhibitor-releasing coupler)

RN 109117-35-3 CAPLUS

CN 1H-Benzotriazolecarboxylic acid, 1-[2-[2-[[2-chloro-5-

[(dodecyloxy)carbonyl]phenyl]amino]-1-(4-methoxybenzoyl)-2-oxoethoxy]-5-hydroxy-4-[(phenylsulfonyl)amino]phenyl]-, phenyl ester (9CI) (CA INDEX NAME)

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(2 CITINGS)

L87 ANSWER 67 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1987:205102 CAPLUS Full-text

DOCUMENT NUMBER: 106:205102

ORIGINAL REFERENCE NO.: 106:33104h,33105a

TITLE: Color photographic coupler

INVENTOR(S): Ichijima, Yasushi; Obayashi, Keiji PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 29 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
JP 61231553	A	19861015	JP 1985-72379	19850405 <	
JP 06090464	В	19941114			
PRIORITY APPLA. INFO.:			JP 1985-72379	19850405 <	

AB A Ag halide color photog. material contains ≥1 compound which releases by coupling reaction with the oxidized form of the principal developer, a precursor of a compound capable of releasing a photog. useful group upon oxidation

IT 108291-77-6

RL: USES (Uses)

(photog. coupler from)

RN 108291-77-6 CAPLUS

CN 1H-Benzotriazolecarboxylic acid, 1-[5-(acetyloxy)-2-[1-[[[5-[[4-[2,4-bis(1,1-dimethylpropyl)phenoxy]-1-oxobutyl]amino]-2-chlorophenyl]amino]carbonyl]-3,3-dimethyl-2-oxobutoxy]-4-ethoxyphenyl]-, phenyl ester (9CI) (CA INDEX NAME)

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___OAc

∽OEt

L87 ANSWER 68 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1987:186316 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 106:186316

ORIGINAL REFERENCE NO.: 106:30057a,30060a

TITLE: Silver halide color photographic material

INVENTOR(S): Ichijima, Yasushi; Yamada, Kozaburo; Obayashi, Keiji

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 29 pp.

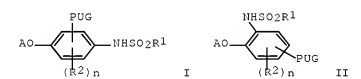
CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
JP 61238057	A	19861023	JP 1985-80021	19850415 <		
PRIORITY APPLN. INFO.:			JP 1985-80021	19850415 <		
GI						



- AB A Ag halide color photog. material contains couplers of the formula I and/or II [A = coupler residue capable of splitting off the group beyond O on reaction with the oxidized form of the developer; R1, R2 = substituent; n = 1-3; PUG = photog. useful group] capable of releasing PUG groups during development.
- IT 108089-14-1

RL: USES (Uses)

(photog. couplers, photog. useful group-releasing)

RN 108089-14-1 CAPLUS

CN 1H-Benzotriazolecarboxylic acid, 1-[2-[2-[[2-chloro-5-[(dodecyloxy)carbonyl]phenyl]amino]-1-(4-methoxybenzoyl)-2-oxoethoxy]-4-methoxy-5-[(phenylsulfonyl)amino]phenyl]-, 4-methoxyphenyl ester (9CI) (CA INDEX NAME)

IT 108107-07-9P

RN

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and use of, as photog. useful group-releasing couplers) 108107-07-9 CAPLUS

CN Pentanamide, N-[5-[[4-[2,4-bis(1,1-dimethylpropyl)phenoxy]-1-

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OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

L87 ANSWER 69 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1987:186305 CAPLUS Full-text

DOCUMENT NUMBER: 106:186305

ORIGINAL REFERENCE NO.: 106:30057a,30060a

TITLE: Silver halide color photographic photosensitive

materials

INVENTOR(S): Ninomiya, Hidetaka; Hirabayashi, Shigeto PATENT ASSIGNEE(S): Konishiroku Photo Industry Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 22 pp.

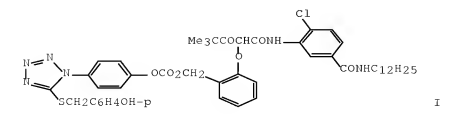
CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
JP 61194443	A	19860828	JP 1985-34506	19850225 <	
JP 05049093	В	19930723			
PRIORITY APPLN. INFO.:			JP 1985-34506	19850225 <	
GI					



AB The claimed color photog. photosensitive materials contain ≥ 1 compound of the formula Y-A-PUG-B (Y = yellow coupler moiety; A = a group separated from

the Y during coupling reaction; B = a protective group for PUG which releases PUG during coupling reaction or by hydrolysis; PUG = photog. useful compound moiety; A can be released together with PUG to give a photog. useful compd of the formula A-PUG. The above PUG-releasing yellow couplers show good chemical stability and good PUG-releasing timing, and hence the color photog. materials show good storage stability, image quality, and sensitivity. Thus, a color photog. paper prepared by using the development

Thus, a color photog. paper prepared by using the development inhibitor-releasing coupler I gave yellow dye images with high Dmax, low fog, and high modulation transfer function.

IT 107758-74-7 107758-76-9 107758-81-6

RL: USES (Uses)

(photog. development inhibitor-releasing yellow coupler)

RN 107758-74-7 CAPLUS

CN Pentanamide, N-[5-[[2-[2,4-bis(1,1-dimethylpropyl)phenoxy]-1-

oxobutyl]amino]-2-chlorophenyl]-2-[[4-[5-[[(4-hydroxyphenoxy)methyl]thio]-1H-tetrazol-1-yl]phenyl]amino]-4,4-dimethyl-3-oxo- (CA INDEX NAME)

RN 107758-76-9 CAPLUS

CN Benzoic acid, 4-chloro-3-[[3-[(4-chlorophenyl)amino]-2-[4-[5-[[(1,3-

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$$\begin{array}{c|c} & & & & \\ & &$$

RN 107758-81-6 CAPLUS
CN Pentanamide, N-[2-chloro-5-[[3-(dodecylsulfonyl)-2-methyl-1-oxopropyl]amino]phenyl]-2-[4-[5-[[(1,1-dioxido-3-oxo-1,2-benzisothiazol-2(3H)-yl)methyl]thio]-1H-tetrazol-1-yl]phenoxy]-4,4-dimethyl-3-oxo-(CAINDEX NAME)

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$$\begin{array}{c|c} & & & \\ & & &$$

L87 ANSWER 70 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1987:1846 CAPLUS $\underline{Full-text}$

DOCUMENT NUMBER: 106:1846
ORIGINAL REFERENCE NO.: 106:371a,374a

TITLE: Use of acylurea compounds for controlling

endoparasites and ectoparasites of warm-blooded

animals

INVENTOR(S): Potter, Michael Fred; Rotramel, George Lorton; Caruso,

Andrew James; Chou, David Teh Wei; Cain, Paul Alfred

PATENT ASSIGNEE(S): Union Carbide Corp., USA SOURCE: PCT Int. Appl., 173 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 8603941	A1	19860717	WO 1985-US2545	19851227 <
M. All BB DK	TH TH	I .TP KR	I.K MW NO SD SH	

	RW:		BE, TD,		CG,	CH,	CM,	DE,	FR,	GA,	GB,	IT,	LU,	ML,	MR,	NL,	SE,	
US	51359	953			Α	1	9920	0804	Ţ	JS 19	85-8	0463	8		19	985120	9 <	<u></u>
AU	86530	006			Α	1	9860	729	P	AU 19	86-5	3006			19	985122	27 <	<u></u>
AU	5993	13			В2		1990	0719										
EP	21100) 4			A1	1	.9870)225	E	EP 19	86-9	0055	3		19	985122	27 <	
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CN	85109	721			Α	1	.9870	715		CN 19	85-1	0972	1		19	985122	28 <	
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FI	86034	490			A	1	.9860	828	E	'I 19	86-3	490			19	986082	28 <	
NO	86034	463			Α	1	.9861	L027	N	10 19	86-3	463			19	986082	28 <	
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PRIORIT	Y APPI	LN, I	NFO.	:					U	IS 19	84-6	8724	9	A	19	84122	8 <	(
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									1	US 19	992-9	2408	39	P	3 19	992080	3 <	.—

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): CASREACT 106:1846; MARPAT 106:1846

The urea derivs. R1CONR2C(Y)NR3R4 [R1 = (un)substituted carbocyclic or heterocyclic ring, etc.; R2, R3 = H, (un)substituted alkyl -benzyl, PhSO2, PhS, etc., R4 = H, R1; Y = O, S] are prepared as endo- and ectoparasiticides. Thus, 3-chloro-4-(4-chloro-1-naphthoxy)-2,5- dimethylaniline (preparation given) was reacted with 2,6-difluorobenzoyl isocyanate in MePh at 50°, to give 1-[3-chloro-4-(4-chloro-1-naphthoxy)-2,5-dimethylphenyl]-3-(2,6-difluorobenzoyl)urea (I). Addition of 25 ppm I to the feed of chicken, totally controlled lice (Menacanthus stramineus).

IT 105621-79-2P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as endo- and ectoparasiticide)

RN 105621-79-2 CAPLUS

CN Benzamide, 2,6-difluoro-N-[[[4-[5-(trifluoromethyl)-1,2,4-oxadiazol-3-yl]phenyl]amino]carbonyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 13 THERE ARE 13 CAPLUS RECORDS THAT CITE THIS RECORD (13 CITINGS)

REFERENCE COUNT: 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 71 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1986:432893 CAPLUS Full-text

DOCUMENT NUMBER: 105:32893

ORIGINAL REFERENCE NO.: 105:5321a,5324a

TITLE: Silver halide photographic materials containing

development inhibitor-releasing photographic couplers

INVENTOR(S): Ono, Mitsunori; Sasaki, Noboru
PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan
SOURCE: Jpn. Kokai Tokkyo Koho, 27 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DAT E	APPLICATION NO.	DATE	
JP 60230139	A	19851115	JP 1984-85835	19840427 <	
JP 05058177	В	19930825			
PRIORITY APPLN. INFO.:			JP 1984-85835	19840427 <	

GI For diagram(s), see printed CA Issue.

AB The claimed Ag halide photog. photosensitive material contains a photog. useful compound-releaser of the formula I (R = coupler moiety; Z = heteroatom which forms an anion when R is released, Z1 = a group of atoms which transport charges toward R1 and forms an electrophilic center; R1 = electron attracting group, atom, or radical; R2 = a photog. useful group; R3 = Z3R4; R4 = nucleophilic group whose reaction with the electrophilic center results in release of R2; Z2, Z3 = bond or a divalent linkage).

IT 102827-64-5

RL: USES (Uses)

(photog. development inhibitor-releasing coupler)

RN 102827-64-5 CAPLUS

CN Pyridinium, 2-[4-[1-[[[2-chloro-5-

[[(dodecyloxy)carbonyl]amino]phenyl]amino]carbonyl]-3,3-dimethyl-2-

oxobutoxy]-2-[(1-ethyl-1H-tetrazol-5-yl)thio]phenyl]-3-hydroxy-1-methyl-,
iodide (1:1) (CA INDEX NAME)

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L87 ANSWER 72 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1986:196909 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 104:196909

ORIGINAL REFERENCE NO.: 104:30989a,30992a

TITLE: Silver halide color photographic photosensitive

materials

INVENTOR(S): Ichijima, Yasushi; Ono, Mitsunori; Sasaki, Noboru

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 27 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 60233649	A	19851120	JP 1984-90437	19840507 <
PRIORITY APPLN. INFO.:			JP 1984-90437	19840507 <

GI For diagram(s), see printed CA Issue.

AB The claimed colorphotog. photosensitive materials contain a compound of the formula I (R = group released during reaction with an oxidized developing

agent; Z = a linkage which releases a group containing Z1 after the releasing of the group R; Z1 = 0, S; A = 4- to 7-membered ring; X = N, O, S; B = b heterocyclic ring having photog. useful compound moiety). the compds. I releases photog. useful compound (such as development inhibitor) with excellent timing, and hence the colorphotog. material exhibit good image sharpness, granularity, and color tone reproducibility. The photog. materials also show excellent storage stability.

IT 102120-55-8P 102120-56-9P

RL: PREP (Preparation)

(preparation and reaction)

RN 102120-55-8 CAPLUS

CN Benzoic acid,

3-[[2-[3-[5-[[[4-(acetyloxy)phenyl]methyl]thio]-1H-tetrazol-1-yl]phenoxy]-4,4-dimethyl-1,3-dioxopentyl]amino]-4-chloro-, hexadecyl ester (CA INDEX NAME)

RN 102120-56-9 CAPLUS

CN Benzoic acid, 4-chloro-3-[[2-[3-(2,5-dihydro-5-thioxo-1H-tetrazol-1-yl)phenoxy]-4,4-dimethyl-1,3-dioxopentyl]amino]-, hexadecyl ester (CA INDEX NAME)

L87 ANSWER 73 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1986:159526 CAPLUS Full-text

DOCUMENT NUMBER: 104:159526

ORIGINAL REFERENCE NO.: 104:25069a,25072a

TITLE: Silver halide color photographic photosensitive

materials

INVENTOR(S): Ichiba, Yasushi; Usui, Hideo; Deguchi, Hisayasu

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 36 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE	
JP 60185950	 A	19850921	JP 1984-33059		19840223	<
JP 05018092	В	19930311				
EP 157146	A2	19851009	EP 1985-101852		19850220	<
EP 157146	A3	19861203				
EP 157146	B1	19890927				
R: CH, DE, FF	R, GB, I	Γ, LI				
US 4618571	A	19861021	US 1985-705473		19850225	<
PRIORITY APPLN. INFO.:			JP 1984-33059	A	19840223	<
ASSIGNMENT HISTORY FOR	US PATE	NT AVAILABLE	IN LSUS DISPLAY FO	DRMAT		
OTHER SOURCE(S):	CASREA	ACT 104:1595	26; MARPAT 104:159	526		
GI						

AΒ Ag halide color photog. photosensitive materials contain a coupler whose coupling reaction with the oxidized developing agent produces a compound which undergoes a redox reaction with a developing agent to release a photog. useful compound The claimed couplers show improved storage stability over that of the couplers with timing groups, and release photog. useful compds. in a controlled manner. The couplers also improve image sharpness, granularity, and color tone reproducibility. Thus, a test photog. film was prepared by using a green-sensitive Ag(Br, I) emulsion containing a magenta coupler 1-(2,4,6-trichlorophenyl)-3-[3-[2-(2,4-di-tertamylphenoxy)butyramido]benzamido-5-oxo-2-pyrazoline (I) and a new magenta coupler II (20 mol% of I). The film was sensitometrically exposed and developed to give Dmax, Dmin, image sharpness (MFT value at 10 cycles/mm), and granularity (RMS value at d. 0.5) of 2.83, 0.13, 114, and 0.0167, resp., vs. 2.95, 0.15, 103, and 0.0210, resp., for a II-free control. ΙT 101124-57-6

RL: USES (Uses)

(photog. development inhibitor-releasing coupler)

RN 101124-57-6 CAPLUS

CN 1H-Benzotriazolecarboxylic acid, 1-[5-[1-[[[2-chloro-5-[(1-oxotetradecyl)amino]phenyl]amino]carbonyl]-3,3-dimethyl-2-oxobutoxy]-2-hydroxy-3-(1,1,3,3-tetramethylbutyl)phenyl]-, phenyl ester (9CI) (CA INDEX NAME)

OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

L87 ANSWER 74 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1986:139248 CAPLUS <u>Full-text</u>

DOCUMENT NUMBER: 104:139248

ORIGINAL REFERENCE NO.: 104:21853a,21856a

TITLE: Colorless ligand-releasing monomers and polymers and

their use to provide dyes with metal ions

INVENTOR(S): Washburn, William N.; Hollister, Kenneth R.

PATENT ASSIGNEE(S): Eastman Kodak Co., USA

SOURCE: U.S., 10 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
	US 4557998	A	19851210	US 1985-688224		19850102 <
	CA 1252463	A1	19890411	CA 1985-482287		19850524 <
	US 4680356	A	19870714	US 1985-764152		19850809 <
	EP 186869	A2	19860709	EP 1985-116273		19851219 <
	EP 186869	A3	19880914			
	EP 186869	B1	19900314			
	R: DE, FR, GB,	NL				
	JP 61166545	A	19860728	JP 1985-293429		19851227 <
PRIOF	RITY APPLN. INFO.:			US 1985-688224	A	19850102 <
ASSIG	SNMENT HISTORY FOR U	S PATEN	T AVAILABLE	IN LSUS DISPLAY F	'ORMAT	
OTHER	R SOURCE(S):	MARPAT	104:139248			
\circ τ						

GΙ

Colorless ligand-releasing polymers are described which can be used as masking dyes for color correction in photog. elements or to form reversal images in photog. elements. The polymer contains (1) recurring units derived from an ethylenically unsatd. polymerizable monomer RCH2:CCoup-Link-Lig (R = H, alkyl; Coup = photog. coupling moiety, Link = coupling group which can be cleaved by an oxidized developer, Lig = a ligand capable of complexing with metal ions. Thus, a 3:1 mol. mixture of a yellow dye-providing color coupler I and N-{{4-chloro-3-{4,4-dimethyl-2-[2,6-di(2-pyridyl)-4-pyridyloxy]-3-oxopentanamido}-phenyl{{-acrylamide-Na2-acrylamido-2-methylpropane-1-sulfonate copolymer (II) was mixed with half their weight of di-Bu phthalate and 3 times their weight with of EtOAc, mixed

their weight of di-Bu phthalate and 3 times their weight with of EtOAc, mixed with gelatin (until homogeneous). The coating levels on a suitable support was gelatin 3.8 g/m2, I 1.8 g/m2, the polymer II 764 mg/m2. The element was imagewise exposed, developed and bleached. No masking dye scale was observed under these conditions but seasoned bleach or dilute ammonium ferrous sulfate solns. generated the magenta color correcting dye scale. 101061-68-1 101061-70-5

IT 101061-68-1 101061

RL: USES (Uses)

(colorless ligand-releasing photog. additive, for application as masking dye for color correction or formation of reversal images)

RN 101061-68-1 CAPLUS

CN 1-Propanesulfonic acid, 2-methyl-2-[(1-oxo-2-propenyl)amino]-, monosodium salt, polymer with N-[2-chloro-5-[(1-oxo-2-propenyl)amino]phenyl]-2-[3,5-di-2-pyridinyl-4-(4-pyridinyl)phenoxy]-4,4-dimethyl-3-oxopentanamide (9CI)

(CA INDEX NAME)

CM 1

CRN 101061-67-0 CMF C37 H32 Cl N5 O4

CM2

CRN 5165-97-9

CMF C7 H13 N O4 S . Na

Na

101061-70-5 CAPLUS RN

1-Propanesulfonic acid, 2-methyl-2-[(1-oxo-2-propenyl)amino]-, monosodium CN salt, polymer with N-[2-chloro-5-[(1-oxo-2-propenyl)amino]phenyl]-4,4dimethyl-3-oxo-2-[4-[6-phenyl-3-(2-pyridinyl)-1,2,4-triazin-5yl]phenoxy]pentanamide (9CI) (CA INDEX NAME)

CM 1

CRN 101061-69-2

CMF C36 H31 C1 N6 O4

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CM 2 CRN 5165-97-9 CMF C7 H13 N O4 S . Na

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OS.CITING REF COUNT: THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD

(4 CITINGS)

REFERENCE COUNT: 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L87 ANSWER 75 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1986:43109 CAPLUS Full-text

DOCUMENT NUMBER: 104:43109 ORIGINAL REFERENCE NO.: 104:6879a,6882a

TITLE: Photographic recording material

INVENTOR(S): Becker, Manfred; Matejec, Reinhart; Endres, Lothar

PATENT ASSIGNEE(S): Agfa-Gevaert A.-G., Fed. Rep. Ger.

SOURCE: Ger. Offen., 43 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
DE 2404054	7.1	10050014	DE 1004 2404054		10040011
DE 3404854	A1	19850814	DE 1984-3404854		19840211 <
EP 152822	A2	19850828	EP 1985-100911		19850130 <
EP 152822	A3	19880127			
EP 152822	B1	19890531			
R: BE, DE, FR,	GB				
US 4636461	A	19870113	US 1985-696901		19850131 <
JP 60258536	A	19851220	JP 1985-22133		19850208 <
JP 05008813	В	19930203			
PRIORITY APPLN. INFO.:			DE 1984-3404854	A	19840211 <
ACCIONMENT HIGHORY HOD I	ישיי אם	NTT 7577 TT 7 DT T	TM TOHO DIODIAN I		

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

MARPAT 104:43109 OTHER SOURCE(S):

AΒ Photog. materials showing high edge effects in the presence of development inhibitor-releasing compds. contain ≥1 iodide-containing Ag halide emulsion in which the grains have zones of different halide compns. The grains are characterized such that from the grain surface to its center there are 3 zones of different halide concentration each following one another and the local iodide concentration is at a maximum at ≥ 1 point that is not on the surface or in the center; the difference between the iodide concentration of the zone with the highest iodide concentration and the iodide concentration of the

furthermost removed zone from the center is preferably ≥ 9 mol%; the portion (in mol% Ag halide) of the zone in which the iodide concentration reaches a maximum is preferably between 20 and 40%, and preferably 70% of the Ag halide grains are cubic or tetradecahedral or in a transition form therebetween. Thus, to a 3-zone gelatin Ag(Br,I) emulsion (AgBr0.995I0.005 30, AgBr0.92I0.08 60, and AgBr 10 mol%) were added a cyan coupler, a DIR compound, and other additives, and the emulsion coated on a support, dried, exposed, and the edge effects at a macrocolor d. of 1.0 for 0.75 g DIR and 1.0 g DIR/AgNO3 determined to be 0.40 and 0.54, resp., vs. 0.27 and 0.25, resp., for a Ag(Br,Cl,I) emulsion.

IT 75956-70-6

RL: USES (Uses)

(photog. development inhibitor-releasing compound, color films with emulsions containing iodide-containing zoned silver halide grains and,

for

improved edge effects)

RN 75956-70-6 CAPLUS

CN Acetamide,

2-(4-chloro-2-tetradecylphenoxy)-N-[4-[4,5-dihydro-5-oxo-4-[(1-

OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD (4 CITINGS)

L87 ANSWER 76 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1985:532313 CAPLUS Full-text

DOCUMENT NUMBER: 103:132313

ORIGINAL REFERENCE NO.: 103:21023a,21026a

TITLE: Color photographic recording material and development

method

INVENTOR(S): Sauerteig, Wolfgang; Ranz, Erwin; Schuetz, Heinz

PATENT ASSIGNEE(S): Agfa-Gevaert A.-G., Fed. Rep. Ger.

SOURCE: Ger. Offen., 33 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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A1
                             19850704
                                        DE 1983-3346621
                                                             19831223 <--
    DE 3346621
    US 4571378
                                       US 1984-680154
                                                              19841210 <--
                      A
                             19860218
    EP 148441
                      A2
                             19850717
                                        EP 1984-115219
                                                              19841212 <--
                      A3
    EP 148441
                             19860625
                             19880217
    EP 148441
                       В1
        R: BE, DE, FR, GB
    JP 60227256
                      A
                             19851112
                                        JP 1984-268710
                                                              19841221 <--
                                                          A 19831223 <--
PRIORITY APPLN. INFO.:
                                        DE 1983-3346621
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OTHER SOURCE(S):
                      MARPAT 103:132313
GΙ
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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The sensitometric characteristics of color photog. materials, especially AΒ color grain in regions of low color d., can be improved, without adversely affecting the sensitivity, by using ≥1 emulsion layer of comparatively higher sensitivity and ≥1 emulsion layer of comparatively lower sensitivity, each having contained therein a coupler-DIR compound combination of different reactivities. The ratio of the effective reaction rate constant of the coupler and the DIR in the higher sensitivity emulsion layer is greater than in the lower sensitivity emulsion layer. Thus, a cellulose acetate support was coated with a red-sensitive gelatin-Aq(Br,I) emulsion layer of lower sensitivity containing I 600, II 30 mg, and a masking coupler, a red-sensitive gelatin-Ag(Br,I) emulsion layer of higher sensitivity containing I 200 and III 40 mg, an interlayer, a green-sensitive layer of lower sensitivity, a green-sensitive layer of higher sensitivity, an interlayer, a yellow filter layer, a blue-sensitive layer of lower sensitivity, a blue-sensitive layer of higher sensitivity, a UV absorber layer, and a top layer. The resultant photog. film was then exposed and developed to show a red sensitivity of 24.1 DIN and a granularity at 0.5 over fog, 1.0 over fog, and 1.5 over fog of 1.5, 1.5, and 1.4, resp., vs. 23.0 DIN, and 1.5, 1.5, and 1.4, resp., for a control containing II in place of III.

IT 75956-70-6

RL: USES (Uses)

(coupler-DIR compound combinations containing, for color photog. films with

improved color grains)

RN 75956-70-6 CAPLUS

CN Acetamide,

2-(4-chloro-2-tetradecylphenoxy)-N-[4-[4,5-dihydro-5-oxo-4-[(1-dihydro-5-0xo-4-[(1-dihydro-5-[(1-dihydro-5-0xo-4-[(1-dihydro-5-[(1-dihydro-5-[(1-dihydro-5-[(1-dihydro-5-[(1-dihydro-5-[

OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

L87 ANSWER 77 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1985:157924 CAPLUS Full-text

DOCUMENT NUMBER: 102:157924

ORIGINAL REFERENCE NO.: 102:24704h,24705a

TITLE: Silver halide color photographic photosensitive

materials

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 27 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patant
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
JP 59206834	A	19841122	JP 1983-82018	19830511 <	
PRIORITY APPLN. INFO.:			JP 1983-82018	19830511 <	
GI					

AB Ag halide color photog. photosensitive materials contain compds. of formula RZZ1R1 (Z = nucleophilic group having N, O, or S as the nucleophilic center or its precursor, Z1 = photog. useful compound moiety; R = group separated from ZZ1R1 during the reaction with oxidized developing agent; R1 = electrophilic group which undergoes intramol. nucleophilic reactor with the

Ι

Z group to form 3- to 7-membered ring after the R-Z bond is broken. The photog. materials having good storage stability and good image quality. Thus, a multilayer color photog. material prepared by using a development inhibitor-releasing coupler I showed improved storage stability and development inhibiting characteristics over those of a control with a conventional development inhibitor-releasing coupler.

IT 95736-67-7

RL: USES (Uses)

(photo. development inhibitor-releasing coupler)

RN 95736-67-7 CAPLUS

OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

L87 ANSWER 78 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1984:510569 CAPLUS Full-text

DOCUMENT NUMBER: 101:110569

ORIGINAL REFERENCE NO.: 101:16869a,16872a

TITLE: Benzoylurea derivatives

PATENT ASSIGNEE(S): Kumiai Chemical Industry Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 59027864	A	19840214	JP 1982-138236	19820809 <
PRIORITY APPLN. INFO.:			JP 1982-138236	19820809 <
GT				

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$$R^1$$
 R^2
 R^2

AB Eight benzoylureas (I, R = H, halo; R1 = halo, NO2; R2 = halo, CF3; X = halo; X1 = CH, N), which had insecticidal activity against Prodenia litura, etc., were prepared either by reaction of 2,6-RXC6H3CONCO (II) with phenylenediamines III or by reaction of 2,6-RXC6H3CONH2 with isocyanatoanilines IV. Thus, treatment of 3 g 50% II (R = X = F) in toluene with 2.4 % III (R1 = NO2, R2 = CF3, X1 = CH) in toluene at room temperature for 1 h gave 84.2% I (R = X = F, R1 = NO2, R2 = CF3, X1 = CH).

II 91590-96-4P 91590-97-5P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation and insecticidal activity of)

RN 91590-96-4 CAPLUS

CN Benzamide, 2-chloro-N-[[[4-(3,5-dichloro-2-pyridinyl)phenyl]amino]carbonyl]- (CA INDEX NAME)

RN 91590-97-5 CAPLUS

CN Benzamide, N-[[[4-(3,5-dichloro-2-pyridinyl)phenyl]amino]carbonyl]-2,6-difluoro- (CA INDEX NAME)

$$\begin{array}{c|c} C1 & & \\$$

L87 ANSWER 79 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1981:165628 CAPLUS Full-text

DOCUMENT NUMBER: 94:165628

ORIGINAL REFERENCE NO.: 94:26927a,26930a

TITLE: Color photographic recording material with a

DIR-coupler of high reactivity

INVENTOR(S): Ranz, Erwin; Lohmann, Joachim; Schuetz, Heinz Dieter

PATENT ASSIGNEE(S): Agfa-Gevaert A.-G., Fed. Rep. Ger.

SOURCE: Ger. Offen., 32 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2853362	A1	19800612	DE 1978-2853362	19781211 <
DE 2853362	C2	19811015		
JP 55084935	A	19800626	JP 1979-8295	19790129 <
US 4315070	A	19820209	US 1980-158990	19800612 <
PRIORITY APPLN. INFO.:			DE 1978-2853362	A 19781211 <
			US 1979-16954	A1 19790302 <

GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB A color photog, recording material contains ≥3 different spectrally sensitized Ag halide emulsion layer units containing nondiffusing dye components and in ≥1 layer a development inhibitor-releasing (DIR) coupler with high reactivity (rate constant >20,000 L mol-1 s-1 at pH 10.2) in a concentration of 10-5-10-3 mol/mol Ag halide or in a concentration of 10-7-10-5 mol/g solid in a Ag halide-free intermediate binder layer adjacent to a Ag halide emulsion layer. Thus, the film tested had the following layers: (1) a red-sensitized Aq(Br, I) emulsion of low sensitivity containing 790 mg cyan coupler, 25 mg DIR coupler I, and 1.6 g gelatin; (2) an intermediate gelatin layer; (3) a green-sensitized Ag(Br,I) emulsion of low sensitivity containing 600 mg magenta coupler, 60 mg DIR coupler II, 80 mg of a masking coupler, and 2 g gelatin; (4) an intermediate gelatin layer; (5) a highly sensitive red-sensitized Ag(Br,I) emulsion containing 250 mg of cyan coupler and 1.0 g gelatin; (6) an intermediate layer containing gelatin, fine-grained AgCl, and DIR coupler III; (7) a highly sensitive green-sensitized Ag(Br,I) emulsion containing 170 mg of magenta coupler, 37

mg of a 2nd magenta coupler, and 2.1 g gelatin; (8) an intermediate gelatin layer; (9) a yellow filter layer; (10) a blue-sensitive layer containing both a sensitive and a relatively unsensitive Ag(Br,I) emulsion with 1.0 g yellow coupler and 2.0 g gelatin; (11) a cover layer of gelatin. The highly reactive DIR coupler IV (k = 50,000 L mol-1 s-1) was added to layer 7 in increasing amts. from $6.07 \times 10-5$ to $3.03 \times 10-4$ mol IV/mol Ag. Upon addition of 1.82 \times 10-4 mol IV/mol Ag the magenta fog was decreased from 0.90 in the IV-free film to 0.83, without change in the sensitivity, gradation, or magenta- and cyan-interimage effects.

IT 75956-70-6

RL: USES (Uses)

(photog. development inhibitor-releasing coupler)

RN 75956-70-6 CAPLUS

CN Acetamide,

 $2-(4-\text{chloro}-2-\text{tetradecylphenoxy})-N-[4-[4,5-\text{dihydro}-5-\text{oxo}-4-[(1-\text{dihydro}-5-\text{dihy$

OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)

L87 ANSWER 80 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1981:123114 CAPLUS Full-text

DOCUMENT NUMBER: 94:123114

ORIGINAL REFERENCE NO.: 94:20143a,20146a

TITLE: Disperse dyes and their use

INVENTOR(S): Neumann, Peter; Elser, Wolfgang; Bock, Gustav; Kermer,

Wolf Dieter

PATENT ASSIGNEE(S): BASF A.-G., Fed. Rep. Ger.

SOURCE: Ger. Offen., 47 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 291242 8	A1	19801009	DE 1979-2912428	19790329 <
US 4373102	A	19830208	US 1980-128156	19800307 <
EP 17132	A1	19801015	EP 1980-101558	19800325 <
EP 17132	B1	19811014		

R: AT, BE, CH, DE, FR, GB, IT, LU, NL, SE

JP 55131064 A 19801011 JP 1980-39143 19800328 <--

JP 63060072 B 19881122

PRIORITY APPLN. INFO.: DE 1979-2912428 A 19790329 <--

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 94:123114

GΙ

$$(NC) 2C$$
 NMe_2
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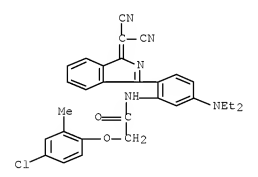
AB Substituted 1-(cyanomethylene)-3-(4-aminophenyl)-1H-isoindole derivs. are prepared and used to dye polyester fibers and polystyrene [9003-53-6] fast blue to violet shades. Thus, 3-(dicyanomethylene)-1-iminoisoindoline [43002-19-3] was heated with N,N-dimethylaniline [121-69-7] in Ac20 containing H2SO4 to give I [76751-73-0], reddish blue on polyester fibers.

IT 76751-35-4

RL: TEM (Technical or engineered material use); USES (Uses) (dye, for polyester fibers, preparation of)

RN 76751-35-4 CAPLUS

CN Acetamide, 2-(4-chloro-2-methylphenoxy)-N-[2-[1-(dicyanomethylene)-1H-isoindol-3-yl]-5-(diethylamino)phenyl]- (CA INDEX NAME)



OS.CITING REF COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS)

L87 ANSWER 81 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1980:613290 CAPLUS Full-text

DOCUMENT NUMBER: 93:213290

ORIGINAL REFERENCE NO.: 93:33906h,33907a

TITLE: Cyan couplers for silver halide color photographic

materials

INVENTOR(S): Kojima, Tamotsu; Fujimatsu, Wataru; Udagawa, Yasushi;

Sasaki, Osamu; Yamashita, Kiyoshi

PATENT ASSIGNEE(S): Konishiroku Photo Industry Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 17 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 55035377	A	19800312	JP 1978-108832	19780904 <
JP 57004896	В	19820128		
PRIORITY APPLN. INFO.:			JP 1978-108832	19780904 <
GT				

$$\begin{bmatrix} \mathbb{R}^2 \text{CONR}^1 \\ \mathbb{H} \text{O} \\ \mathbb{F} \end{bmatrix} \mathbb{R}^2 \mathbb{R}^2$$

AB Ag halide color photog. materials contain cyan couplers of the formula I [R = H, C1-5 aliphatic hydrocarbon moiety with/without substituent; R1 = H, organic moiety; R2 = diffusion-resistant moiety conventionally used in color couplers; R1R2 in combination may complete N-containing heterocyclic ring; Z = O-containing organic moiety which is bonded via O to the active position of the coupler moiety; Z1 = simple bond, or n-valent organic moiety, or H (when n = 1); n = 1,2]. Thus, a cyan coupler

 $6-[\alpha-(2,4-\text{di-tert-amylphenoxy})\,\text{butyramido}]-4-\text{ethoxycarbonylmethoxy-}2-fluoro-3-methylphenol was used to give a color photog. material, which gave photog. images with good stability, optical d., and high color-formation speed even in the absence of PhCH2OH in a color developer solution$

IT 75505-50-9P

RN 75505-50-9 CAPLUS

CN Ethanedioic acid, 1-[5-[[2-[2-(2H-benzotriazol-2-yl)-4-(1,1-dimethylpropyl)phenoxy]acetyl]amino]-3-fluoro-4-hydroxy-2-methylphenyl] 2-ethyl ester (CA INDEX NAME)

L87 ANSWER 82 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1979:195580 CAPLUS Full-text

DOCUMENT NUMBER: 90:195580

ORIGINAL REFERENCE NO.: 90:30969a,30972a

TITLE: Couplers for silver halide color photographic

materials

INVENTOR(S): Ishikawa, Hitoshi; Fujiwara, Mitsuhito; Kikuchi,

Shoji; Wada, Hajime; Endo, Takanari

PATENT ASSIGNEE(S): Konishiroku Photo Industry Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 28 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 53141622	A	19781209	JP 1977-56190	19770516 <
JP 60039218	В	19850905		
PRIORITY APPLN. INFO.:			JP 1977-56190	19770516 <

GI For diagram(s), see printed CA Issue.

AΒ Ag halide color photog. materials contain ≥1 coupler selected form compds. of the general formulas I, II, and III (R, R1, R2, R3, R4, R5, R6, R7 = H, OH, NH2, NO2, halo, alkyl, alkenyl, aryl, alkoxy, alkenyloxy, aryloxy, acyloxy, alkylthio, alkenylthio, arylthio, monoalkylamino, dialkylamino, acylamino, sulfonamido, carbamoyl, sulfamoyl, sulfo, sulfino, R9 Zm; RR1, R1R2, or R2R3 combination may form a 5- or 6-membered ring; R8 = R9Zm, alkyl, alkenyl, cycloalkyl, aryl, acyl, alkylsulfonyl, arylsulfonyl, carbamoyl, sulfamoyl, oxalyl, oxamoyl, oxycarbonyl, oxalacetyl groups having substituents selected from alkyl, aryl and alkoxy group; R9 = cyan; magentaor yellow-coupler moiety; Z = an organic moiety; m = 0, 1; $n \ge 1$, Z1 = a divalent moiety obtained by removing a H atom from R8]. The dye images obtained from the above couplers exhibit excellent light fastness and moisture resistance. Thus, a multilayer color photog. paper was prepared by using the yellow coupler IV, the magenta coupler V, and the cyan coupler VI. The color photog. paper was imagewise exposed and developed to give a color image which showed good light and moisture resistances.

IT 69964-06-3

RL: TEM (Technical or engineered material use); USES (Uses) (photog. cyan coupler)

RN 69964-06-3 CAPLUS

CN Acetamide, 2-[2-(2H-benzotriazol-2-yl)-4-(1,1-dimethylethyl)phenoxy]-N-

(3,5-dichloro-2-hydroxy-4-methylphenyl)- (CA INDEX NAME)

$$\begin{array}{c} \text{Me} \\ \text{Cl} \\ \text{NH-C-CH}_2 - \text{O} \\ \text{N} \\ \text{N} \\ \text{N} \end{array}$$

IT 69964-17-6

RL: TEM (Technical or engineered material use); USES (Uses) (photog. magenta coupler)

RN 69964-17-6 CAPLUS

CN Acetamide,

N-[4-chloro-3-[[4,5-dihydro-5-oxo-1-(2,4,6-trichlorophenyl)-1Hpyrazol-3-yl]amino]phenyl]-2-[4-(1,1-dimethylethyl)-2-[5[(methylsulfonyl)amino]-2H-benzotriazol-2-yl]phenoxy]- (CA INDEX NAME)

IT 69964-24-5

RL: TEM (Technical or engineered material use); USES (Uses) (photog. yellow coupler)

RN 69964-24-5 CAPLUS

CN Pentanamide, N-[5-[[2-[2-(2H-benzotriazol-2-yl)-4-(1,1-dimethylethyl)phenoxy]acetyl]amino]-2-chlorophenyl]-4,4-dimethyl-3-oxo-(CA INDEX NAME)

L87 ANSWER 83 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1979:14636 CAPLUS Full-text

DOCUMENT NUMBER: 90:14636

ORIGINAL REFERENCE NO.: 90:2323a,2326a

TITLE: Color complexes with moieties absorbing ultraviolet

rays

INVENTOR(S): Van Poucke, Raphael Karel; Vanden Eynde, Hector

Alfons; Monbaliu, Marcel Jacob

PATENT ASSIGNEE(S): Agfa-Gevaert N. V., Belg.

SOURCE: Fr. Demande, 30 pp.

CODEN: FRXXBL

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
FR 2363133	A1	19780324	FR 1977-25623		19770822 <
FR 2363133	B1	19800711			
PRIORITY APPLN, INFO.:			GB 1977-19578	A	19770510 <
GI					

Obtained with the use of color formers, which are chemical bonded to a UV-absorbing moiety, i.e. in the form of I (Z = divalent organic groups; R1, R2 = substituted or unsubstituted alkyl and aryl; and R = color former radical). Thus, a red-sensitized photog. Ag(Br,I) emulsion (2.3 mol% I-) containing 0.006 mol II and 0.012 mol 2,2-dimethyl-6-hydroxy-7-isononylchroman per 0.032 mol Ag halide was coated on a suitable support, dried, imagewise exposed, developed in a N,N-diethyl-p-phenylenediamine type developer, and treated with a bleach-fix bath containing K3Fe(CN)6 and Na2S2O3 to give a cyan image which showed no change in color after 15-h exposure to radiation from a 1500-W Xe lamp.

IT 68658-25-3

RL: USES (Uses)

(photog. color coupler, for high-stability color image production)

RN 68658-25-3 CAPLUS

CN Butanamide, 2-[4-[4,6-bis(2,4-dimethylphenyl)-1,3,5-triazin-2-yl]-3-hydroxyphenoxy]-N-[4-chloro-3-[[4,5-dihydro-5-oxo-1-(2,4,6-trichlorophenyl)-1H-pyrazol-3-yl]amino]phenyl]- (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

AUTHOR(S):

CORPORATE SOURCE:

L87 ANSWER 84 OF 84 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1973:487360 CAPLUS Full-text

DOCUMENT NUMBER: 79:87360

ORIGINAL REFERENCE NO.: 79:14122h,14123a

TITLE: Irreversible enzyme inhibitors. 198.

Diaminodihydro-s-triazines and diaminopyrimidines bearing substituted (ureidomethyl)phenyl substituents as reversible inhibitors of dihydrofolate reductase Ashton, Wallace T.; Kirk, Larry L.; Baker, B. R. Dep. Chem., Univ. California, Santa Barbara, CA, USA

SOURCE: Journal of Medicinal Chemistry (1973), 16(5), 453-6

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE: Journal LANGUAGE: English

Triazines bearing m-(arylureidomethyl)phenyl substituents and pyrimidines bearing p-(arylureidomethyl)phenyl substituents were inhibitors of dihydrofolate reductase [9002-03-3] from L1210 mouse leukemia cells, and of cultured L1210 cells. The most potent compound, 2,4-diamino-6-methyl-5-[p-[(3-nitrophenyl)ureidomethyl]phenyl]pyrimidine (I) [41935-22-2], inhibited L1210 cells at 61 pM in vitro and was thus 2000 times as active as 4,6-diamino-1,2-dihydro-2,2-dimethyl-1-[m-(m-fluorosulfonylphenylureidomethyl)phenyl]-s-triazine ethanesulfonate [19159-37-6], a compound with antileukemic activity in vivo. The enhanced activity of I and some other compds. on the cells was probably due to improved membrane transport, since all the compds. caused similar inhibition of the enzyme. Activity was correlated with the electron-withdrawing power of the

meta substituent on the terminal benzene ring. To synthesize I, 5-(p-aminopheny1)-2, 4-diamino-6-methylpyrimidine [41935-24-4] was diazotized and reacted with CuCN to give the 5-(p-cyanopheny1) compound, which was reduced with H2/Pt to the 5-(p-aminomethyl)phenyl compound and reacted with Ph N-(3-nitrophenyl)carbamate (prepared from Ph chloroformate [1885-14-9] and m-nitroaniline) to yield I.

IT 50699-45-1P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, dihydrofolate reductase and leukemia inhibiting activities

by)

RN 50699-45-1 CAPLUS

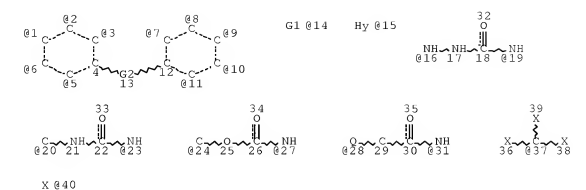
CN Urea, N-(3-chlorophenyl)-N'-[[4-(2,4-diamino-6-methyl-5-pyrimidinyl)phenyl]methyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD (4 CITINGS)

FILE 'HOME' ENTERED AT 14:19:25 ON 14 SEP 2011

SEARCH HISTORY

=> d stat que 141; d his nofile L10 STR



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VAR G2=16-4 19-12/16-12 19-4/20-4 23-12/20-12 23-4/24-4 27-12/24-12 27-4/

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VPA 15-1/2/3/5/6 U

VPA 14-7/8/9/10/11 U

NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM

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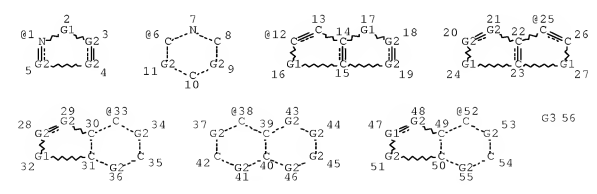
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GRAPH ATTRIBUTES:

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NUMBER OF NODES IS 40

STEREO ATTRIBUTES: NONE L11 STR



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VAR G3=1/6/12/25/33/38/52

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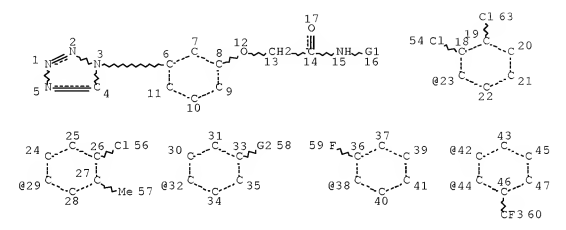
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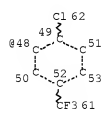
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L34 STF



Page 1-A



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D SCA SEL RN

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             1 SEA SPE=ON ABB=ON L70 AND L71
             7 SEA SPE=ON ABB=ON (L73 OR L74 OR L75 OR L76 OR L77 OR L78)
1 SEA SPE=ON ABB=ON (L47 OR L48 OR L49 OR L50 OR L51 OR L52 OR
L79
L80
                L53 OR L54) AND L41
L81
              1 SEA SPE=ON ABB=ON (L47 OR L48 OR L49 OR L50 OR L51 OR L52 OR
                L53 OR L54) AND L18
              7 SEA SPE=ON ABB=ON (L79 OR L81)
L82
     FILE 'CAPLUS' ENTERED AT 14:17:02 ON 14 SEP 2011
                D QUE NOS L82
                D IBIB ABS HITSTR L82 1-7
     FILE 'REGISTRY' ENTERED AT 14:17:24 ON 14 SEP 2011
                D STAT QUE L41
     FILE 'CAPLUS' ENTERED AT 14:17:33 ON 14 SEP 2011
                D OUE NOS L42
L83
            164 SEA SPE=ON ABB=ON L42 NOT L82
            144 SEA SPE=ON ABB=ON L83 AND PATENT/DT
L84
              7 SEA SPE=ON ABB=ON (L83 NOT L84) AND PY<2004
L85
             77 SEA SPE=ON ABB=ON L84 AND (PD<20030829 OR AD<20030829 OR
L86
                PRD<20030829)
             84 SEA SPE=ON ABB=ON (L85 OR L86)
L87
                D IBIB ABS HITSTR L87 1-84
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FILE 'HOME' ENTERED AT 14:19:25 ON 14 SEP 2011 D STAT QUE L41